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Scientific and Technical Information Center SCIENTIFICATION SCIENTIFIC Angell Examiner #: \$8697 Date: \$17/02 AUDIT 1635 Phone Number 30-605: U.S. Serial Number: \$94740.574 Fill Normal Bildy Room Location #11212. CMI Results Formal Preferred (circle): PAPER DISK E-MAIL OU. 12015: CMI more than one searCRF submitted, please prioritize searches in order of need: more than one searCRF submitted, please prioritize searches in order of need: **The search of the learn to fine tearch topic, and describe as specifically as possible the subject matter to be searched. The search topic, and describe as specifically as possible the subject matter to be searched. The search one search of the learn to fine tearch topic, and describe as specifically as possible the subject matter to be searched. The search of the cover sheet, pertinent claims, and abstract. Fille of Invention: Define anyloses, appecial meaning. Give examples or relevant citations, authors, ethnif nown. Please attach a copy of the cover sheet, pertinent claims, and abstract. Fille of Invention: DNA Vaccine. For farm animals, in particular hovines and parcines inventors (please provide full names): Smoon Rossun Le Roux Smoon Rossun Le Roux Smoon Rossun Le Roux Earliest Priority Filling Date: 3/20/2000 For Squance Searches Only Please include all pertinent information (parent; child, divisional, or issued patent numbers) along with the appropriate provide of the control of the structure of the specific parent child, divisional, or issued patent numbers) along with the appropriate provide of the specific parent child, divisional, or issued patent numbers) along with the appropriate parent in unbers. All Claims are affected for your convenience. Only Claims 1:11, 13-21, 44-55, One of the control of the specific parent pare
quester's Full Name: Jon Eric Angell Examiner #: 38697 Date: 8/15/07: t Unit: 1535 Phone Number 30-605-1165 Serial Number: 0017-005794 ail Box and Bldg/Room Location 11217- CMI Results Format Preferred (circle): PAPER DISK E-MAIL olice 12015- CMI more than one search 15 submitted, please prioritize searches in order of need. more than one search 15 submitted, please prioritize searches in order of need. more than one search 15 submitted, please prioritize searches in order of need. more than one search 15 submitted, please prioritize searches in order of need. more than one search 15 submitted, please prioritize searches in order of need. more than one search 15 submitted, please prioritize search 15 in order of need. more than one search 15 submitted, please prioritize search 15 in order of need. more than one search 15 submitted, please prioritize search 15 in order of need. more than one search 15 submitted, please prioritize search 15 in order of need. more than one search 15 submitted, please include a special meaning. Give examples or relevant citations, authors, etc. of the inventors of the inventors of the inventors, and abstract 15 in order 16 in order 17 in order 1
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se provide a detailed statement of the search topic, and describe as specifically as possible the subject maker to be searched, see provide a detailed statement of the search topic, and describe as specifically as possible the subject maker to be searched, see provide the concept or side the elected species or structures, keywords, synonyms, acronyms, and registry numbers, and combine-with the concept or yof the invention. Define any terms that may have a special meaning. Give examples or relevant citations, authors, etcylif with Please attach a copy of the cover sheet, pertinent claims, and abstract. The variety of the cover sheet, pertinent claims, and abstract. Smora Barzy-Le-Roux Thiest Priority Filling Date: 3/20/2000 or Sequence Searches Only* Please include all pertinent information (parent, child, divisional, or issued patent numbers) along with the propriate artist number. The structure of the structure Searchy for the structure of acclaims. All Claims are affected for your convenience. Only claims 1-11, 18-21, 44-55. All Claims are affected for your convenience. Only claims 1-11, 18-21, 44-55. The structure of the formula of claim 1 issued for all the propriate and purposes. The structure of the formula of claim 1 issued for all the purposes. The structure of the formula of claim 1 issued for all the purposes. The structure of the formula of claim 1 issued for all the purposes. The structure of the formula of claim 1 issued for all the purposes. The structure of the formula of claim 1 issued for all the purposes. The formula of the formula of claim 1 issued for all the purposes. The formula of the formula of claim 2 issued for all the purposes. The formula of the formula of claim 2 issued for all the purposes. The formula of the formula of the formula of claim 2 issued for all the purposes. The formula of
de the elected species of students that may have a special meaning. Give examples or relevant citations, administration, volumes and particular forms and structure. The sea attach a copy of the cover sheet, pertinent claims, and abstract. The of Invention: DNA Vaccine for form animals, in particular forwing and parcines are controlled for a superintent information (parent, child, divisional, or issued patent numbers) along with the corpulatorial number. The office a structure Search for this, divisional, or issued patent numbers) along with the reprintental number. The office a structure search for your convenience. Only claims 1-11, 18-21, 44-55. All Claims are affected for your convenience. Only claims 1-11, 18-21, 44-55. The structure of the familia of claim 1 is used for different purposes. The following and different for course pigs for your cantendary for the purposes. The following and different for course pigs for your claims for any faither classifications, Journal Technical Info Specials. The chain all not specials. Edward Hart Technical Info Specials. STIC/Biolechia.
le of Invention: DNA Vaccine for farm animals, in particular formes and parcines rentors (please provide full names): Jean-Christophe Andonnet, Lawent Fischer, Simona Barzu-Le-Roux Thiest Priority Filing Date: 3/20/2000 or sequence Searches Only* Please include all pertinent information (parent, child, divisional, or issued patent numbers) along with the propriageerial number. I would like a structure search for the structure of claim! The structure of the formula of claim 1 is used for different purposes. DNB / huckers and of interfet to course if pigs for vaccination purposes. DNB / huckers and of interfet to course if pigs for vaccination purposes. Please days height to Contact me for any further classification / quidance Please days height to Contact me for any further classification / quidance Technical Info Specialist STIC Blotech 1980 1980 1980 1980 1980 1980 1980 1980
entors (please provide full names): Simona Barzu-Le-Roux Thiest Priority Filing Date: 3/20/2000 or Sequence Searches Only* Please include all pertinent informations (parent, child, divisional, or issued patent numbers) along with the ropriate serial number. I would like a structure search for the structure of Claim 1. The structure of the formula of claim 1 is used for antivery and 60-63 are pending in the application. The structure of the formula of claim 1 is used for antivery.
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FILE COVERS 1907 - 16 Aug 2002 VOL 137 ISS 8 FILE LAST UPDATED: 15 Aug 2002 (20020815/ED)

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GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 14

STEREO ATTRIBUTES: NONE

112 SEA FILE=REGISTRY SSS FUL L3 L5

STR L7

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S 6
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REP G1=(11-17) C VAR G2=OH/N REP G3=(2-3) C NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 14

STEREO ATTRIBUTES: NONE

L9 109 SEA FILE=REGISTRY SUB=L5 SSS FUL L7
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L11 ANSWER 1 OF 32 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 2002:10302 HCAPLUS

DOCUMENT NUMBER: 136:74555

TITLE: Vaccine against foot-and-mouth disease

INVENTOR(S): King, Andrew; Burman, Alison; Audonnet,

Jean-Christophe; Lombard, Michel

PATENT ASSIGNEE(S): Merial, Fr.

SOURCE: PCT Int. Appl., 79 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: French

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	PATENT NO.				KIND DATE				APPLICATION NO'. DATE									
WO.	2002	00029	5.1	Δ.	1	20020	0103		W	200	01-FI	R2042	2 :	2001	0627			
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		CM	UР	нп	TD	TT.	TN.	TS.	JP.	KE,	KG,	KΡ,	KK,	KΖ,	ا بالل	rv'	LK,	
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		RO.	RU.	SD.	SE.	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	
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	144.	DE.	DK.	ES.	FI,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	ΝL,	PT,	SE,	TR,	BF,	
		BJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	TG			

FR 2000-8437 20000629 20020104 FR 2810888 Α1 20010627 AU 2001-70678 AU 2001070678 Α5 20020108 20000629 FR 2000-8437 A PRIORITY APPLN. INFO.: 20010627 W WO 2001-FR2042

MARPAT 136:74555 OTHER SOURCE(S):

The invention concerns a vaccine against foot-and-mouth disease, using as antigen an efficient amt. of empty capsids of the foot-and-mouth virus, said empty capsids being obtained by expressing, in eukaryotic cells, cDNA of the Pl region of the foot-and-mouth virus genome coding for the capsid and cDNA of the region of the foot-and-mouth virus genome coding for protease 3C, the vaccine further comprising a carrier or excipient pharmaceutically acceptable in veterinary medicine. invention also concerns the insertion of a mutation in the sequence VP2 (introducing a cysteine), thereby stabilizing the empty capsids and the resulting viruses.

153312-64-2, Dmrie TΤ

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(vaccine against foot-and-mouth disease)

THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS 7 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT REFERENCE COUNT:

L11 ANSWER 2 OF 32 HCAPLUS COPYRIGHT 2002 ACS

2001:886529 HCAPLUS ACCESSION NUMBER:

136:32635 DOCUMENT NUMBER:

Improved methods of transfection of cells with a TITLE:

receptor targeted vector and uses thereof

Hart, Stephen Lewis INVENTOR(S):

Ich Productions Ltd., UK PATENT ASSIGNEE(S): PCT Int. Appl., 111 pp. SOURCE:

CODEN: PIXXD2

Patent DOCUMENT TYPE: English LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATI	ENT 1	NO.		KIND DATE				APPLICATION NO. DATE									
WO :	W:	CO, GM, LS, RO, UZ, GH, DE, BJ,	AG, CR, HR, LT, RU, VN, GM, DK, CF,	CU, HU, LU, SD, YU, KE, ES, CG,	AM, CZ, ID, LV, SE, ZA, LS,	DE, IL, MA, SG, ZW, MW, FR.	AU, DK, IN, MD, SI, AM, MZ, GB,	DM, IS, MG, SK, AZ, SD, GR, GN,	BA, DZ, JP, MK, SL, SL, IE, GW,	BB, EC, KE, MN, TJ, KG, SZ, IT, ML,	BG, EE, KG, MW, TM, KZ, TZ, LU, MR,	KP, MX, TR, MD, UG, MC, NE,	BY, FI, KR, MZ, TT, RU, ZW, NL, SN, A	2001(BZ, GB, KZ, NO, TZ, TJ, AT, PT, TD, 2000	CA, GD, LC, NZ, UA, TM BE, SE, TG	LK, PL, UG, CH, TR,	LR, PT, US,
									US 2	001-	2874	10P	Ρ	2001	0501		coll

The present invention relates to an improved method of transfecting cells. Transfection of confluent cells or other slowly dividing or non-dividing AB cells that are in contact with each other with a nucleic acid using a non-viral receptor targeted vector may be improved by the concurrent use of an agent that disrupts cell-cell junctions, esp. EGTA. The vector is esp. an integrin-targeting transfection vector complex comprising (i) a nucleic acid, esp. a nucleic acid encoding a sequence of interest, (ii) an integrin-binding component, esp. an integrin-targeting peptide, (iii) a polycationic nucleic acid-binding component, esp. an oligolysine, and (iv)

a lipid component, esp., DOPE, DOTMA, DOSPA or combinations thereof. Various applications of the improved method of transfection are described. 168479-03-6, DOSPA

RL: ARG (Analytical reagent use); BSU (Biological study, unclassified); ANST (Analytical study); BIOL (Biological study); USES (Uses) (improved methods of transfection of cells with a receptor targeted vector and uses thereof)

L11 ANSWER 3 OF 32 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2001:886528 HCAPLUS

DOCUMENT NUMBER:

136:32634

TITLE:

IT

Integrin-binding peptides and their use in increasing the efficiency of transformation of animal cells in

vector vaccines for cancer, respiratory and

heart diseases

INVENTOR(S):

Hart, Stephen Lewis
Ich Productions Ltd., UK

PATENT ASSIGNEE(S): SOURCE:

PCT Int. Appl., 108 pp.

CODEN: PIXXD2

DOCUMENT TYPE: LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

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PATENT NO. KIND DATE APPLICATION NO. DATE

WO 2001092542 A2 20011206 WO 2001-GB2394 20010530

W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

PRIORITY APPLN. INFO::

GB 2000-13090 A 20000530

US 2001-287410P P 20010501
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A method of increasing the efficiency of transformation of animal cells by binding the transforming the DNA to integrins is described. Peptides contg. an integrin-binding motif and a polylysine sequence for binding nucleic acid are used to bring the DNA in close contact with the cell. The peptide-nucleic acid complex may be delivered in a liposome. The nucleic acid preferably is or relates to a gene that is the target for gene therapy, gene vaccination or antisense therapy. The integrin binding component comprises an integrin-binding element and a spacer element. integrin binding element is an integrin binding peptide and contains a cyclic conserved RGD amino acid sequence. The spacer element is a peptide that is longer and/or more hydrophobic than the dipeptide spacers GG (glycine-glycine) and GA (glycine-alanine), contains an .epsilon.-amino hexanoic acid, is the the N terminus of the integrin-binding element and has enhanced transfection activity. The lipid component preferably has membrane destabilizing or fusogenic properties like DOPE, DOTMA, DOSPA or combinations thereof. An embodiment of the present invention provides a ratio of lipid component (DOPE or DOTMA): integrin-binding/polycationic nucleic acid-binding component: nucleic acid of 0.75:4:1 by wt. or 0.5:1.25:0.25 nmol. Furthermore, the present invention provides a ratio of lipid component (DOPE or DOSPA): integrin-binding/polycationic nucleic acid-binding component: nucleic acid of 12:4:1 by wt. Transfection of

confluent cells or other slowly dividing or non-dividing cells that are in contact with each other with a nucleic acid using a non-viral receptor targeted vector may be improved by the concurrent use of an agent that disrupts cell-cell junctions, like the calcium chelator EGTA (at concns. of less than 1 mM) or an antibody like anti-cadherin. The present invention can be used in a vaccines for neuroblastoma, leukemias and other cancers as well as for diseases affecting smooth muscle and cardiac muscle tissues as well as for respiratory dieases. These vectors are also useful as a kit for improved transfection activity and they can deliver very large DNA mols. to cells.

158571-62-1, Lipofectamine 168479-03-6, DOSPA RL: BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(liposomes for nucleic acid delivery contg.; integrin-binding peptides and their use in increasing efficiency of transformation of animal cells in vector vaccines for cancer, respiratory and heart diseases)

L11 ANSWER 4 OF 32 HCAPLUS COPYRIGHT 2002 ACS 2001:798084 HCAPLUS

ACCESSION NUMBER: 135:348865

Compositions and methods for in vivo delivery of DOCUMENT NUMBER: TITLE:

polynucleotide-based therapeutics

Hartikka, Jukka; Sukhu, Loretta; Manthorpe, Marston

INVENTOR(S): Vical Incorporated, USA PATENT ASSIGNEE(S): PCT Int. Appl., 176 pp. SOURCE:

CODEN: PIXXD2

Patent DOCUMENT TYPE: English LANGUAGE:

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

APPLICATION NO. DATE KIND DATE PATENT NO. _____ ____ WO 2001-US12975 20010423 20011101 A2 WO 2001080897

RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, W: CA, JP, US

PT, SE, TR US 2001-839574 20010423 20020214 US 2000-198823P P 20000421 US 2002019358 PRIORITY APPLN. INFO.: US 2000-253153P P 20001128

The present invention relates to pharmaceutical compns. and methods to improve expression of exogenous polypeptides into vertebrate cells in AB vivo, utilizing delivery of polynucleotides encoding such polypeptides. More particularly, the present invention provides the use of salts, in particular sodium and potassium salts of phosphate, in aq. soln., and auxiliary agents, in particular detergents and surfactants, in pharmaceutical compns. and methods useful for direct polynucleotide-based polypeptide delivery into the cells of vertebrates.

153312-64-2, Dmrie 208040-06-6, Gap dlrie

RL: MOA (Modifier or additive use); PEP (Physical, engineering or chemical process); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)

(compns. and methods for in vivo delivery of polynucleotide-based therapeutics)

L11 ANSWER 5 OF 32 HCAPLUS COPYRIGHT 2002 ACS 2001:791879 HCAPLUS ACCESSION NUMBER:

135:335117 DOCUMEN'T NUMBER:

TITLE:

Immunological adjuvants containing Hemagglutinating virus-containing charged liposomes, and manufacture

APPLICATION NO. DATE

thereof

INVENTOR(S):

Honda, Kazuo; Kaneda, Yasushi; Shiozaki, Koichi Chemo-Sero-Therapeutic Research Institute, Japan

PATENT ASSIGNEE(S):

Jpn. Kokai Tokkyo Koho, 9 pp.

SOURCE:

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.	KIND	DATE	APPLICATION NO.	
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IT

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (charged liposomes contg. Hemagglutinating virus and lipids as immunol. adjuvants)

L11 ANSWER 6 OF 32 HCAPLUS COPYRIGHT 2002 ACS 2001:545519 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

135:142202

TITLE:

Improved DNA vaccines for livestock

INVENTOR(S):

Audonnet, Jean-Christophe Francis; Fischer, Laurent

Bernard; Barzu-le-Roux, Simona

PATENT ASSIGNEE(S):

SOURCE:

Merial, Fr. PCT Int. Appl., 79 pp.

CODEN: PIXXD2

DOCUMENT TYPE: LANGUAGE:

Patent French

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

		_	KIND	DATE		AP	PLIC	CATIC	N NC).	DATE			
	TENT NO		KIND			 W.C	200)1-FF	 R187		20010)119		
WO FF US	RW: 28040 20020	52888 AE, AG, CR, CU, HU, ID, LU, LV, SD, SE, ZA, ZW, GH, GM, DE, DK, BJ, CF,	A3 AL, Al CZ, D. IL, I MA, M SG, S AM, A KE, L ES, F CG, C A1 A1	20010726 20011220 M, AT, AU, E, DK, DM, N, IS, JP, D, MG, MK, I, SK, SL, Z, BY, KG, S, MW, MZ, T, FR, GB, 1, CM, GA, 20010727 20020516	AZ, DZ, KE, MN, TJ, KZ, SD, GR,	BA, EE, KG, MW, TM, MD, SL, IE, GW,	BB, ES, KP, MX, TR, RU, SZ, IT, ML, R 20 S 20	BG, FI, KR, MZ, TT, TZ, LU, MR, 00-7	BR, GB, KZ, NO, TZ, TM UG, MC, NE, 98	BY, GD, LC, NZ, UA, XW, NL, SN,	LK, PL, UG, AT, PT, TD, 2000 2001	LR, PT, UZ, BE, SE, TG 0121 0116	LS, RO, VN, CH, TR,	LT, RU, YU,

MARPAT 135:142202

OTHER SOURCE(S): The invention concerns a DNA vaccine against a pathogen affecting livestock, in particular cattle and swine, comprising a plasmid contg. a nucleotide sequence coding for an immunogen of a pathogen of the animal species concerned, in conditions enabling the expression in vivo of said sequence, and a cationic lipid contg. a quaternary ammonium salt, of formula R1-O-CH2-CH(OR1)-CH2-N+(CH3)2-R2 X-, wherein: R1 is a linear aliph. radical, satd. or unsatd., having 12 to 18 carbon atoms; R2 is another aliph. radical, contg. 2 or 3 carbon atoms; and X is a hydroxyl or amine group, said lipid being preferably DMRIE.

153312-64-2, Dmrie IT

RL: PEP (Physical, engineering or chemical process); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses) (improved DNA vaccines for livestock)

L11 ANSWER 7 OF 32 HCAPLUS COPYRIGHT 2002 ACS 2001:490587 HCAPLUS

ACCESSION NUMBER:

DOCUMENT NUMBER:

135:362424

TITLE:

Highly efficient gene delivery by mRNA electroporation in human hematopoietic cells: superiority to

lipofection and passive pulsing of mRNA and to electroporation of plasmid cDNA for tumor antigen

loading of dendritic cells

AUTHOR(S):

Van Tendeloo, Viggo F. I.; Ponsaerts, Peter; Lardon, Filip; Nijs, Griet; Lenjou, Marc; Van Broeckhoven, Christine; Van Bockstaele, Dirk R.; Berneman, Zwi N. Laboratory of Experimental Hematology, Antwerp

CORPORATE SOURCE:

University Hospital, University of Antwerp, Antwerp,

Belg.

SOURCE:

Blood (2001), 98(1), 49-56 CODEN: BLOOAW; ISSN: 0006-4971 American Society of Hematology

PUBLISHER: DOCUMENT TYPE:

Journal

English

Designing effective strategies to load human dendritic cells (DCs) with LANGUAGE: tumor antigens is a challenging approach for DC-based tumor AB vaccines. Here, a cytoplasmic expression system based on mRNA electroporation to efficiently introduce tumor antigens into DCs is described. Preliminary expts. in K562 cells using an enhanced green fluorescent protein (EGFP) reporter gene revealed that mRNA electroporation as compared with plasmid DNA electroporation showed a markedly improved transfection efficiency (89% vs. 40% EGFP+ cells, resp.) and induced a strikingly lower cell toxicity (15% death rate with mRNA vs. 51% with plasmid DNA). Next, mRNA elec. troporation was applied for nonviral transfection of different types of human DCs, including monocyte-derived DCs (Mo-DCs), CD34+ progenitor-derived DCs (34-DCs) and Langerhans cells (34-LCs). High-level transgene expression by mRNA electroporation was obtained in more than 50% of all DC types. MRNA-electroporated DCs retained their phenotype and maturational potential. Importantly, DCs electroporated with mRNA-encoding Melan-A strongly activated a Melan-A-specific cytotoxic T lymphocyte (CTL) clone in an HLA-restricted manner and were superior to mRNA-lipofected or -pulsed DCs. Optimal stimulation of the CTL occurred when Mo-DCs underwent maturation following mRNA transfection. Strikingly, a nonspecific stimulation of CTL was obsd. when DCs were transfected with plasmid DNA. The data clearly demonstrate that Mo-DCs electroporated with mRNA efficiently present functional antigenic peptides to cytotoxic T cells. Therefore, electroporation of mRNA-encoding tumor antigens is a powerful technique to charge human dendritic cells with tumor antigens and could serve applications in future DC-based tumor vaccines.

189203-05-2, DMRIE-C ΙT

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(lipofection with; highly efficient gene delivery by mRNA electroporation in human hematopoietic cells for tumor antigen loading

of dendritic cells)

THERE ARE 32 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT REFERENCE COUNT:

L11 ANSWER 8 OF 32 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2001:409275 HCAPLUS

DOCUMENT NUMBER:

136:198465

TITLE:

Vaxfectin enhances antigen specific antibody titers and maintains Th1 type immune responses to plasmid DNA

immunization

AUTHOR(S):

Reyes, L.; Hartikka, J.; Bozoukova, V.; Sukhu, L.; Nishioka, W.; Singh, G.; Ferrari, M.; Enas, J.;

Wheeler, C. J.; Manthorpe, M.; Wloch, M. K.

CORPORATE SOURCE:

Department of Cell Biology, Vical Incorporated, San

Diego, CA, 92121, USA

SOURCE:

Vaccine (2001), 19(27), 3778-3786 CODEN: VACCDE; ISSN: 0264-410X

Elsevier Science Ltd. PUBLISHER:

DOCUMENT TYPE:

Journal English

Antigen specific immune responses were characterized after i.m. LANGUAGE: immunization of BALB/c mice with 5 antigen encoding plasmid DNAs (pDNAs) complexed with Vaxfectin, a cationic lipid formulation. Vaxfectin increased IgG titers for all of the antigens with no effect on the CTL responses to the 2 antigens for which CTL assays were performed. Both antigen specific IgG1 and IgG2a were increased, although IgG2a remained greater than IgG1. Furthermore, Vaxfectin had no effect on IFN-.gamma. or IL-4 prodn. by splenocytes re-stimulated with antigen, suggesting that the Th1 type responses typical of i.m. pDNA immunization were not altered. Studies with IL-6 -/- mice suggest that the antibody enhancement is IL-6 dependent and results in a correlative increase in antigen specific antibody secreting cells.

370108-99-9, Vaxfectin TΤ

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL

(Biological study); USES (Uses) (Vaxfectin enhanced antigen-specific antibody titers maintaining Th1

type immune responses to plasmid DNA vaccines)

THERE ARE 56 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT 56 REFERENCE COUNT:

L11 ANSWER 9 OF 32 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2001:168152 HCAPLUS

DOCUMENT NUMBER:

134:221435

TITLE:

Prevention of myocarditis, abortion and intrauterine infection associated with porcine circovirus-2

INVENTOR(S):

Ellis, John Albert; Allan, Gordon Moore; Meehan, Brian; Clark, Edward; Haines, Deborah; Hassard, Lori;

Harding, John; Charreyre, Catherine Elisabeth; Chappuis, Gilles Emile; Krakowka, George Steve; Audonnet, Jean-Christophe Francis; McNeilly, Francis

PATENT ASSIGNEE(S):

Merial, Fr.; University of Saskatchewan; The Queen's

University of Belfast PCT Int. Appl., 133 pp.

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

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APPLICATION NO. DATE
                   KIND DATE
    PATENT NO.
                                      _____
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                                     WO 2000-EP8781
                                                      20000828
                         20010308
                   A2
    WO 2001016330
       W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU,
           MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI,
           SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
       RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
           DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ,
           CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                                      20000828
                                       BR 2000-14155
                         20020507
    BR 2000014155
                   Α
                                    US 1999-151564P P 19990831
PRIORITY APPLN. INFO .:
                                                   A 20000531
                                    US 2000-583350
                                                    W 20000828
                                    WO 2000-EP8781
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The invention is based on the discovery that porcine circovirus (PCV-2) is a causative agent of myocarditis, abortion and intrauterine infection, as AB well as post-weaning multisystemic wasting syndrome in pigs. Thus, immunol. compns. contg. the recombinant poxvirus for inducing an immunol. response in aa host animal to which the immunol. compn. is administered. Also described are methods of treating or preventing disease caused by PCV-2 by administering the immunol. compns. of the invention to an animal in need of treatment or susceptible to infection by PCV-2. Such immunol. compns. comprise (1) attenuated or inactivated strains of PCV-2, (2) plasmid vectors expressing open reading frames of PCV-2 and vaccination of pigs with DNA formulated with DMRIE, DMRIE-DOPE, or carbomer adjuvants, and (3) a recombinant poxvirus, such as the canarypox virus (Rentschler strain) contg. foreign DNA encoding the major capsid virus or ORF1 or ORF2 from PCV-2.

153312-64-2, DMRIE IT

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (adjuvant; prevention of myocarditis, abortion and intrauterine infection assocd. with porcine circovirus-2)

L11 ANSWER 10 OF 32 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2001:167832 HCAPLUS

DOCUMENT NUMBER:

134:212748

TITLE:

Lipid-nucleic acid compositions for stimulating cytokine secretion and inducing an immune response Semple, Sean C.; Harasym, Troy O.; Klimuk, Sandra K.;

INVENTOR(S):

Kojic, Ljiljiana D.; Bramson, Jonathan L.; Mui,

Barbara; Hope, Michael J.

PATENT ASSIGNEE(S):

Inex Pharmaceuticals Corp., Can. PCT Int. Appl., 94 pp.

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

 TENT 2001	 26	KII 	 DATE 			_					DATE 2000			
2001	26		 2001 AT, DK,	70 F T	70.77	BA, EE,	BB, ES,	BG, FI,	BR, GB,	BY, GD,	BZ, GE,	CA, GH,	CH, GM,	CN, HR,

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HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
             LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             SD, SE, SG
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ,
             CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                                                20000828
                                             BR 2000-13834
                             20020423
    BR 2000013834
                        Α
                                             EP 2000-956004
                                                                20000828
             AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL
                             20020612
    EP 1212085
                                          US 2000-176406P P
                                                               20000113
PRIORITY APPLN. INFO.:
                                                            W 20000828
                                          WO 2000-CA1013
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Lipid-nucleic acid particles can provide therapeutic benefits, even when the nucleic acid is not complementary to coding sequences in target cells. It has been found that lipid-nucleic acid particles, including those contg. non-sequence specific oligodeoxynucleotides, can be used to stimulate cytokine secretion, thus enhancing the overall immune response of a treated mammal. Further, immune response to specific target antigens can be induced by administration of an antigenic mol. in assocn. with lipid particles contg. non-sequence specific oligodeoxynucleotides. The nucleic acid which is included in the lipid-nucleic acid particle can be a phosphodiester (i.e., an oligodeoxynucleotide consisting of nucleotide residues joined by phosphodiester linkages) or a modified nucleic acid which includes phosphorothicate or other modified linkages, and may suitably be one which is non-complementary to the human genome, such that it acts to provide immunostimulation in a manner which is independent of conventional base-pairing interactions between the nucleic acid and nucleic acids of the treated mammal. In particular, the nucleic acid may suitably contain an immune-stimulating motif such as a CpG motif, or an immune stimulating palindromic sequence. The cationic lipid included in the nucleic acid particles may be suitably selected from among DODAP, DODMA, DMDMA, DOTAP, DC-Chol, DDAB, DODAC, DMRIE, DOSPA and DOGS. In addn., the lipid particle may suitably contain a modified aggregation-limiting lipid such as a PEG-lipid, a PAO-lipid or a ganglioside.

RL: PEP (Physical, engineering or chemical process); THU (Therapeutic 168479-03-6, DOSPA ΙT use); BIOL (Biological study); PROC (Process); USES (Uses) (DOSPA; lipid-nucleic acid compns. for stimulating cytokine secretion and inducing an immune response)

153312-64-2, DMRIE RL: PEP (Physical, engineering or chemical process); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses) (lipid-nucleic acid compns. for stimulating cytokine secretion and inducing an immune response)

L11 ANSWER 11 OF 32 HCAPLUS COPYRIGHT 2002 ACS 2001:146642 HCAPLUS ACCESSION NUMBER:

135:330213

ΙT

Vaxfectin enhances the humoral immune response to DOCUMENT NUMBER: TITLE:

plasmid DNA-encoded antigens

Hartikka, J.; Bozoukova, V.; Ferrari, M.; Sukhu, L.; AUTHOR(S):

Enas, J.; Sawdey, M.; Wloch, M. K.; Tonsky, K.;

Norman, J.; Manthorpe, M.; Wheeler, C. J.

Department of Cell Biology, Vical Incorporated, San CORPORATE SOURCE:

Diego, CA, 92121, USA

Vaccine (2001), 19(15-16), 1911-1923 SOURCE:

CODEN: VACCDE; ISSN: 0264-410X

Elsevier Science Ltd. PUBLISHER:

Journal DOCUMENT TYPE:

This report characterizes Vaxfectin, a novel cationic and neutral lipid LANGUAGE: formulation which enhances antibody responses when complexed with an antigen-encoding plasmid DNA (pDNA). In mice, i.m. injection of Vaxfectin formulated with pDNA encoding influenza nucleoprotein (NP) increased antibody titers .ltoreq. 20-fold, to levels that could not be reached with pDNA alone. As little as 1 .mu.g of pDNA formulated with Vaxfectin per muscle resulted in higher anti-NP titers than that obtained with 25 .mu.g naked pDNA. The antibody titers in animals injected with Vaxfectin-pDNA remained higher than in the naked pDNA controls for at least 9 mo. The enhancement in antibody titers was dependent on the Vaxfectin dose and was accomplished without diminishing the strong anti-NP cytolytic T cell response typical of pDNA-based vaccines. In rabbits, complexing pDNA with Vaxfectin enhanced antibody titers .ltoreq. 50-fold with needle and syringe injections and also augmented humoral responses when combined with a needle-free injection device. Vaxfectin did not facilitate transfection and/or increase synthesis of .beta.-galactosidase reporter protein in muscle tissue. ELISPOT assays performed on bone marrow cells from vaccinated mice showed that Vaxfectin produced a 3- to 5-fold increase in the no. of NP-specific plasma cells. Thus, Vaxfectin should be a useful adjuvant for enhancing pDNA-based vaccinations.

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(Vaxfectin enhances the humoral immune response to plasmid DNA-encoded antigens)

ΙT

RL: PRP (Properties); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(Vaxfectin enhances the humoral immune response to plasmid DNA-encoded

antigens)

AUTHOR(S):

THERE ARE 53 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT 53 REFERENCE COUNT:

L11 ANSWER 12 OF 32 HCAPLUS COPYRIGHT 2002 ACS 2001:121937 HCAPLUS

ACCESSION NUMBER:

135:225548 DOCUMENT NUMBER:

Effects of different transfection reagents on genetic TITLE:

immunization of rabies virus glycoprotein cDNA Zhang, Mao-lin; Hu, Rong-liang; Yu, Xing-long; Tu,

Chang-chun; Qian, Ai-dong; Rong, Ai-hong; Li,

Hong-wei; Yin, Zhen

The Militry Veterinary Institute, Quartermaster CORPORATE SOURCE:

University of PLA, Changchun, 130062, Peop. Rep. China

Zhongguo Shouyi Xuebao (2000), 20(6), 528-531

SOURCE: CODEN: ZSXUF5; ISSN: 1005-4545

Zhongguo Shouyi Xuebao Bianjibu

PUBLISHER: Journal DOCUMENT TYPE:

Three rabies virus glycoprotein expressing vectors, pGFP-C1-RGP, pSV2-RGP LANGUAGE: and pcDNA3-RGP were constructed by cloning rabies virus glycoprotein cDNA AΒ into pGFP-C1, pSV2-dhfr and pcDNA3, resp. Expression of all three vectors was confirmed on cells and in newborn mouse brains. The highest expression level was achieved when the rabies virus glycoprotein gene was regulated by CMV promoter/enhancer. After 3 times of inoculations at intervals of 2 wk in the form of naked DNA, DNA-lipofectamine and DNA-PEI complex, specific antibodies against rabies virus were detected in sera of mice by means of ELISA. The antibody titer went up with the increase of the amt. of plasmids injected. However, when the amt. of the plasmid went beyond 20 .mu.g/mouse, there was no pos. correlation between the dose of DNA injected and the level of immune response when PEI and lipofectamine were used as transfection reagents. The plasmid vaccination could protect mice from the challenge of CVS. Long lasting humoral immune responses were proved with ELISA and PCR amplification 6 mo after the primary inoculation.

158571-62-1, Lipofectamine ΙT

RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(effects of different transfection reagents on genetic immunization of rabies virus glycoprotein cDNA)

L11 ANSWER 13 OF 32 HCAPLUS COPYRIGHT 2002 ACS 2001:101291 HCAPLUS

ACCESSION NUMBER:

134:161880 DOCUMENT NUMBER:

cDNAs encoding the Flt-3 receptor ligand and there use TITLE:

as adjuvants in vector vaccines

Hermanson, Gary George INVENTOR(S):

Vical Inc., USA PATENT ASSIGNEE(S):

PCT Int. Appl., 148 pp. SOURCE:

CODEN: PIXXD2

Patent DOCUMENT TYPE: English LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2001009303	A2	20010208	WO 2000-US20679	20000731
WO 2001009303	A3	20010816		

W: CA, JP, US

RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,

PT, SE PRIORITY APPLN. INFO.:

US 1999-146170P P 19990730

A method of increasing the strength of the immune response of vector vaccines using an expression vector for the Flt3 ligand is described. The vaccines are made of independent non-integrating expression vectors: one encodes the antigen or a cytokine and the other encodes the Flt3 ligand. The present invention also provides a method broadly directed to improving immune response of a vertebrate in need of immunotherapy by administering in vivo, into a tissue of a vertebrate, a Flt-3 ligand-encoding polynucleotide and one or more antigen- or cytokine-encoding polynucleotides. The polynucleotides are incorporated into the cells of the vertebrate in vivo, and a prophylactically or therapeutically effective amt. of a Flt-3 ligand and one or more antigens is produced in vivo.

153312-64-2, DMRIE 208040-06-6, GAP-DLRIE

299207-54-8, GAP-DMORIE

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (in delivery of vector vaccines; cDNAs encoding Flt-3 receptor ligand and there use as adjuvants in vector vaccines

L11 ANSWER 14 OF 32 HCAPLUS COPYRIGHT 2002 ACS

2001:64121 HCAPLUS ACCESSION NUMBER:

134:136654

DOCUMENT NUMBER:

Feline calicivirus genes and vaccines, in TITLE:

particular recombined vaccines

Audonnet, Jean-Christophe Francis; Baudu, Philippe Guy INVENTOR(S):

Nicolas; Brunet, Sylvie Claudine

PATENT ASSIGNEE(S):

Merial, Fr.

SOURCE:

PCT Int. Appl., 61 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

French

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAS	TENT 1	, O		KIND DATE				A!	PPLIC	CATIO	ON NO). 					
WO	2001	0059	34	A2	2	2001	0125		W	O 200	00-FI	3205	1	2000	0713		
WO		AE, CR, HU, LU, SD, YU,	AG, CU, ID, LV, SE, ZA,	AL, CZ, IL, MA, SG, ZW,	AM, DE, IN, MD, SI, AM,	AT, DK, IS, MG, SK, AZ,	AU, DM, JP, MK, SL, BY,	MN, TJ, KG,	KG, MW, TM, KZ,	KP, MX, TR, MD,	KR, MZ, TT, RU,	KZ, NO, TZ, TJ, UG,	LC, NZ, UA, TM	LK, PL, UG,	LR, PT, US, BE,	LS, RO, UZ,	LT, RU, VN,
		DE,	DK,	ES,	FI,	FR,	GB,	GR, GW.	ΙΕ,	MR,	NE,	SN,	TD,	TG	00,	D.,	ВJ,
FR AU	2796 2796 2000 2000 2000 1228 R:	396 397 0657 0125 193 AT,	65 12 BE,	A A A A CH,	1 5 .2 DE	2001 2001 2001	0119 0119 0205 0402 0807 ES,	FR,	F A B E GB,	R 19 R 20 U 20 R 20 CP 20 GR,	99-9 00-1 00-6 00-1 00-9	761 5765 2512 5324 LI,	3 LU,	2000 2000 2000 2000 2000 NL,	0713 0713 0713 MC,	IE,	SI,
PRIORIT	Y APE				110		,		FR 2	999- 2000- 2000-	1761		Α	1999 2000 2000	0211		

MARPAT 134:136654 OTHER SOURCE(S):

The invention concerns the sequence of the capsid gene and a corresponding cDNA sequence, of a dominant FCV strain called FCV 431. The invention also concerns the capsid gene sequence and the cDNA sequence of a complementary strain called G1. The cDNA sequences can be incorporated in expression vectors for prepg. immunogenic formulations and recombined vaccines or subunits providing vaccination against the feline calicivirus disease.

153312-64-2, Dmrie ΙT

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); PEP (Physical, engineering or chemical process); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses) (adjuvant; feline calicivirus genes and vaccines)

L11 ANSWER 15 OF 32 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2000:900790 HCAPLUS

DOCUMENT NUMBER:

134:55493

TITLE:

Porcine circovirus vaccine

INVENTOR(S):

Audonnet, Jean-christophe Francis; Bublot, Michel; Perez, Jennifer Maria; Charreyre, Catherine Elisabeth

Merial, Fr.

PATENT ASSIGNEE(S): SOURCE:

PCT Int. Appl., 40 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

FAMILY ACC. NUM. COUNT:

English

PATENT INFORMATION:

PATENT NO.

KIND DATE

APPLICATION NO. DATE

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                                               WO 2000-EP5611 20000608
                               20001221
                       A2
    WO 2000077188
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR,
                               20010531
    WO 2000077188
             LV, MA, MD, MG, MA, MN, MM, MA, NO, NZ, FL, FI, KO, KO, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
             AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
     EP 1185659
              IE, SI, LT, LV, FI, RO
                                                                      20000608
                                                  BR 2000-11733
                        A 20020723
                                              US 1999-138352P P 19990610
     BR 2000011733
                                                                  W 20000608
PRIORITY APPLN. INFO.:
                                              WO 2000-EP5611
                             MARPAT 134:55493
     The invention relates to immunogenic prepns. or vaccines
OTHER SOURCE(S):
     comprising, on the one hand, a plasmid vector encoding and expressing a
     gene from porcine circovirus (PCV), in particular selected from the group
      consisting of ORF1 of PCV-2, ORF2 of PCV-2, ORF1 of PCV-1 and ORF2 of
      PCV-1, and , on the other hand, an element capable of increasing the
      immune response directed against the product of expression of the gene,
      which can be a carbomer, a porcine cytokine, e.g. GM-CSF or a cationic
      lipid of formula (I), in which R1 is a satd. or unsatd. linear aliph.
      radical having from 12 to 18 carbon atoms, R2 is another aliph. radical comprising from 2 to 3 carbon atoms, and X is a hydroxyle or amine group.
      The cationic lipid can be DMRIE, possibly coupled with DOPE.
      Vaccines contg. plasmid vector encoding and expressing a gene from
      porcine circovirus were prepd. and tested against PMWS.
      RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
      153312-64-2, DMRIE
 ΙT
          (vaccine comprising, cationic lipid or neutral lipid; porcine
          circovirus vaccine)
 L11 ANSWER 16 OF 32 HCAPLUS COPYRIGHT 2002 ACS
                              2000:900679 HCAPLUS
 ACCESSION NUMBER:
                              134:55491
                              DNA vaccines against Paramyxoviridae for
 DOCUMENT NUMBER:
                              pets and game animals and their delivery in liposomes
 TITLE:
                               containing cationic lipids
                               Fischer, Laurent Jean-Charles; Barzu-le, Roux Simona;
                               Audonnet, Jean-Christophe Francis
  INVENTOR(S):
                               Merial, Fr.
  PATENT ASSIGNEE(S):
                               PCT Int. Appl., 110 pp.
  SOURCE:
                               CODEN: PIXXD2
                               Patent
  DOCUMENT TYPE:
                               French
  LANGUAGE:
  FAMILY ACC. NUM. COUNT:
  PATENT INFORMATION:
                                                  APPLICATION NO. DATE
                          KIND DATE
        PATENT NO.
                                  _____
                          ____
                                                                         20000608
                                                   WO 2000-FR1592
                                   20001221
        WO 2000077043
                           A2
                AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU,
        WO 2000077043
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ANGELL 09 / 760574
             LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD,
         SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ,
             CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                                                19990610
                                             FR 1999-7604
                             20001215
                       A1
     FR 2794648
                                                                20000608
                                              BR 2000-11732
                        Α
                              20020305
     BR 2000011732
                                                                20000608
                                             EP 2000-940474
                             20020313
                        A2
     EP 1185662
             AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO
                                                             A 19990610
                                           FR 1999-7604
PRIORITY APPLN. INFO.:
                                           US 1999-144490P P 19990719
                                                             W 20000608
                                           WO 2000-FR1592
                          MARPAT 134:55491
OTHER SOURCE(S):
     The invention aims at improving the efficacy and protection induced by DNA
     vaccination against viruses of the family of Paramyxoviridae and against
     the herpes virus, in pets and sport animals. The improvement of DNA
     vaccination is achieved either by formulating the vaccine with a
     cationic lipid contg. a quaternary ammonium salt, DMRIE, or by
     modifications in the nucleotide sequence coding for the antigen of
     interest in particular of deletions of the fragment of the nucleotide
     sequence coding for the transmembrane domain of the antigen of interest,
     and/or insertions of introns and/or insertions of nucleotide sequences
     coding for the signal peptides, or by adding GM-CSF, or by combinations
     thereof. The invention also concerns the resulting vaccines.
```

completely protected a group of five dogs challenged with the virus. 153312-64-2, DMRIE ΙT

RL: MOA (Modifier or additive use); THU (Therapeutic use); BIOL

(Biological study); USES (Uses) (in liposomes for delivery of DNA vaccines; DNA vaccines against Paramyxoviridae for pets and game animals and

their delivery in liposomes contg. cationic lipids)

series of expression vectors for antigen genes of canine distemper virus and felid, canid, and equid herpes viruses that used the signal sequence of a tissue plasminogen activator gene were constructed by std. methods. In some cases, derivs. lacking the transmembrane domain were used to improve secretion of the extracellular domain. Expression vectors also carrying the genes for cytokines, esp. colony-stimulating factor 2 were also constructed. Use of genes for colony-stimulating factor 2 derived from the target host is demonstrated. A combination of vectors carrying genes for the fusion protein and hemagglutinin of canine distemper virus

L11 ANSWER 17 OF 32 HCAPLUS COPYRIGHT 2002 ACS 2000:861646 HCAPLUS

ACCESSION NUMBER:

134:21482

DOCUMENT NUMBER: TITLE:

INVENTOR(S):

Cytofectin dimers and methods of use thereof

Wheeler, Carl J. Vical, Inc., USA

PATENT ASSIGNEE(S): SOURCE:

PCT Int. Appl., 50 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2000073263 WO 2000073263	A1 C2	20001207 20020711	WO 2000-US14676	20000526

W: CA, JP, US

RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,

PT, SE

EP 2000-939373 20000526 20020306 Α1 EP 1183231

AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,

IE, FI

PRIORITY APPLN. INFO.:

US 1999-136472P P 19990528 WO 2000-US14676 W 20000526

OTHER SOURCE(S):

MARPAT 134:21482

GΙ

Me CH2O(CH2)13Me Me(CH2)13OCH2 Me $\texttt{Me}\,(\texttt{CH}_2)\, \texttt{13} \texttt{OCHCH}_2 \texttt{N}\,(\texttt{CH}_2)\, \texttt{3} \texttt{CONHCHCONH}\,(\texttt{CH}_2)\, \texttt{3} \texttt{NCH}_2 \texttt{C}\, \texttt{HO}\,(\texttt{CH}_2)\, \texttt{13} \texttt{Me}$ CH₂ Me

I

A compn. is provided comprising a novel cationic lipid compd. having AΒ hydrophobic tails and two quaternary ammonium headgroups bridged by a linker. The compn. is useful as a cytofectin for facilitating delivery and transfection of biol. active agents, particularly anionic bioactive agents such as DNA, into cells. The compn. is useful also as an adjuvant for enhancing the humoral immune response of a vertebrate to an immunogen, esp. an immunogen encoded by a polynucleotide-based vaccine. In certain preferred embodiments, the cationic lipid compd. is a dimer contg. quaternary ammonium headgroups bridged by a linker having DNA and/or cell receptor binding affinity, such as a polypeptide or polyamine. Also disclosed is an immunogenic compn. comprising an immunogen and the compn. of the present invention. I was prepd. as an example compd.

310445-42-2P 310445-43-3P 310445-44-4P RL: BPR (Biological process); BSU (Biological study, unclassified); SPN ΙT (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses)

(cationic lipids prepn. as cytofectin for delivery and transfection of biol. agents)

153312-64-2, Dmrie ΙT

RL: RCT (Reactant); RACT (Reactant or reagent) (cationic lipids prepn. as cytofectin for delivery and transfection of biol. agents)

282533-25-9P IT

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(cationic lipids prepn. as cytofectin for delivery and transfection of

biol. agents) REFERENCE COUNT:

THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 18 OF 32 HCAPLUS COPYRIGHT 2002 ACS 2000:850412 HCAPLUS

ACCESSION NUMBER: DOCUMENT NUMBER:

134:365419

TITLE:

Large-scale feasibility of gene transduction into

human CD34+ cell-derived dendritic cells by

adenoviral/polycation complex

Di Nicola, Massimo; Carlo-Stella, Carmelo; Milanesi, Marco; Magni, Michele; Longoni, Paolo; Mortarini, AUTHOR(S):

Roberta; Anichini, Andrea; Tomanin, Rosella; Scarpa,

Maurizio; Gianni, A. Massimo

Division of Medical Oncology, Istituto Nazionale

Tumori, Milan, 20133, Italy

CORPORATE SOURCE: British Journal of Haematology (2000), 111(1), 344-350 SOURCE:

CODEN: BJHEAL; ISSN: 0007-1048

Blackwell Science Ltd.

PUBLISHER: Journal DOCUMENT TYPE:

With a view to using multiple injections of anticancer dendritic cell LANGUAGE:

(DC)-based vaccines, we evaluated the feasibility of the adenoviral transduction of large amts. of human CD34+ cell-derived DCs, and analyzed the persistence of the transgene expression and the integrity of DC functional activity after the transduction/cryopreservation procedures. Mature DCs generated from highly enriched human CD34+ cells were transduced by a recombinant adenovirus (rAd-MFG) that carried a modified, membrane-exposed, alk. phosphatase (AP) sequence as the reporter gene. Cationic lipids such as LipofectAmine or poly-L-lysine were mixed with the viral particles before the transduction of the target cells. highest transduction efficiency was obtained at a multiplicity of infection (MOI) rate of 500 (AP + DCs: 50 .+-. 2%, viability = 95%) under both small- and large-scale conditions. The addn. of poly-L-lysine or LipofectAmine increased the percentage of transduced cells at an MOI of 500 (CDla+/AP+ cells = 85 .+-. 3% and 80 .+-. 2% resp.). Polycations made it possible to reduce the amts. of viral particles, with high efficiency of transduction being achieved at a MOI of 100 with 10 .mu.g/mL poly-L-lysine (CD1a+/AP+: 68 .+-. 9%) or 30 .mu.g/mL LipofectAmine (CD1a+/AP+: 60 .+-. 7%). Evaluation of the immunophenotype of the transduced DCs showed that the lack of a DC subpopulation was more susceptible to adenoviral transduction. Cryopreservation of transduced DCs did not modify the viability or percentage of AP+ cells that maintain antigen-presenting cell (APC) functions. These findings indicate the efficacy of this method for the transduction of large amts. of CD34+ cell-derived DCs using small quantities of adenoviral vector mixed with polycations. Cryopreservation of transduced DCs did not damage their viability or APC functions, thus making it possible to plan multiple injections of engineered DC-based vaccines.

IT

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study)

(large-scale feasibility of gene transduction into human CD34+ cell-derived dendritic cells by adenoviral/polycation complex)

THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT 26 REFERENCE COUNT:

L11 ANSWER 19 OF 32 HCAPLUS COPYRIGHT 2002 ACS 2000:707018 HCAPLUS

ACCESSION NUMBER:

Adjuvant compositions and methods for enhancing immune DOCUMENT NUMBER: TITLE:

responses to polynucleotide-based vaccines

Wheeler, Carl J.

INVENTOR(S): Vical Incorporated, USA PATENT ASSIGNEE(S): PCT Int. Appl., 72 pp.

SOURCE: CODEN: PIXXD2

Patent DOCUMENT TYPE: English LANGUAGE:

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

APPLICATION NO. DATE KIND DATE PATENT NO. -----____ WO 2000-US8282 20000324 20001005 A2 WO 2000057917 20010104 WO 2000057917 AЗ RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE 20000324 EP 2000-919777 20020102 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, EP 1165140 IE, FI US 1999-126340P P 19990326 W 20000324 WO 2000-US8282

PRIORITY APPLN. INFO.:

The invention provides adjuvants, immunogenic compns., and methods useful for polynucleotide-based vaccination and immune response. In particular, the invention provides an adjuvant of cytofectin: co-lipid mixt. wherein cytofectin is GAP-DMORIE.

153312-60-8, DORIE 153312-64-2, DMRIE IT 154486-25-6, GAP-DMRIE 188949-12-4, DMORIE 199171-54-5, DLRIE 208040-06-6, GAP-DLRIE 282533-23-7, DOSPA 299207-54-8, GAP-DMORIE

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); PEP (Physical, engineering or chemical process); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses) (adjuvant compns. contg. cytofectin:co-lipid mixts. and methods for enhancing immune responses to polynucleotide-based vaccines)

L11 ANSWER 20 OF 32 HCAPLUS COPYRIGHT 2002 ACS 2000:580894 HCAPLUS ACCESSION NUMBER:

133:155214

AUTHOR(S):

LipofectAMINE coated hepatitis C virus core gene DOCUMENT NUMBER: vaccine promotes efficacy of immune responses TITLE:

Feng, Zhihua; Zhou, Yongxing; Wang, Quanchu; Du, Dewei; Jiao, Chengsong; Li, Jinge; Li, Guangyu Department of Infectious Diseases, Tangdu Hospital,

Fourth Military Medical University, Xi an, 710038, CORPORATE SOURCE:

Peop. Rep. China

Disi Junyi Daxue Xuebao (2000), 21(7), 817-819 SOURCE:

CODEN: DJDXEG; ISSN: 1000-2790 Disi Junyi Daxue Xuebao Bianjibu

PUBLISHER: Journal DOCUMENT TYPE: Chinese

The efficacy of LipofectAMINE coated recombinant plasmid-contg. hepatitis LANGUAGE: C virus (HCV) core gene inductive immune responses was studied. The HCV core gene coding region was inserted into the eukaryotic expression plasmid pcDNA3, and then the recombinant plasmid pcDNAHCV-C was constructed and expressed transiently with LipofectAMINE in the SP2/0 cells. After purifn., these plasmids directly or encapsuled with LipofectAMINE were injected into BALB/c mice. HCV core antibody from immunized mice was detected by ELISA. The enzyme-cutting identification showed that HCV core gene fragment was cloned into pcDNA3 eukaryote vectors. HCV C antibody was pos. in sera of 12 mice immunized and was time-dependent. The HCV C antibody titer for core antigen induced by plasmid encapsulated with lipofectamine was higher than of control mice. The results showed that liposome technique combined with gene vaccine can promote the efficacy of immune responses in BALB/c mice.

158571-62-1, Lipofectamine ΙT

ANGELL 09 / 760574

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study)

(humoral immune response to genetic immunization with hepatitis C core antigen is enhanced by LipofectAMINE encapsulation of plasmid vector)

L11 ANSWER 21 OF 32 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2000:573482 HCAPLUS

DOCUMENT NUMBER:

134:146025

TITLE:

Effectiveness of combined interleukin 2 and B7.1 vaccination strategy is dependent on the sequence and order: A liposome-mediated gene therapy treatment for

bladder cancer

AUTHOR(S):

Larchian, William A.; Horiguchi, Yutaka; Nair, Smita K.; Fair, William R.; Heston, Warren D. W.; Gilboa,

Eli

CORPORATE SOURCE:

Department of Urology, The Cleveland Clinic

Foundation, Cleveland, OH, 44195, USA

SOURCE:

Clinical Cancer Research (2000), 6(7), 2913-2920

CODEN: CCREF4; ISSN: 1078-0432

PUBLISHER:

American Association for Cancer Research

DOCUMENT TYPE: Journal LANGUAGE: English

The authors have developed a novel liposome-mediated immunogene therapy using interleukin 2 (IL-2) and B7.1 in a murine bladder cancer model. A carcinogen-induced murine bladder cancer cell line, MBT-2, was transfected with cationic liposome 1,2-dimyristyloxypropyl-3-dimethyl-hydroxyethyl ammonium bromide/dioleolylphosphatidylethanolamine and IL-2 plasmid. The optimized transfection condition generated IL-2 levels of 245-305 ng/106 cells/24 h, 100-fold higher than the levels seen with retrovirus transfection. Ninety percent of the peak level of IL-2 prodn. was maintained for up to 11 days after transfection. Animal studies were conducted in C3H/HeJ female mice with 2.times.104 MBT-2 cells implanted orthotopically on day 0. Multiple vaccination schedules were performed with i.p. injection of 5.times.106 IL-2 and/or B7.1 gene-modified cell prepns. The greatest impact on survival was obsd. with the day 5, 10, and 15 regimen. Control animals receiving retrovirally gene-modified MBT-2/IL-2 cell prepns. had a median survival of 29 days. Animals receiving the IL-2 liposomally gene-modified cell prepn. alone had a median survival of 46 days. Seventy-five percent of animals receiving IL-2 followed by B7.1 gene-modified tumor vaccines were the only group to show complete tumor-free survival at day 60. All of these surviving animals rejected the parental MBT-2 tumor rechallenge and survived at day 120 with a high CTL response. Thus, liposome-mediated transfection demonstrates a clear advantage as compared with the retroviral system in the MBT-2 model. Multi-agent as opposed to single-agent cytokine gene-modified tumor vaccines were beneficial. These "targeted" sequential vaccinations using IL-2 followed by B7.1 gene-modified tumor cells increased a systemic immune response that translated into increased survival.

IT 153312-64-2, DMRIE

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(liposome contg.; combined interleukin 2 and B7.1 vaccination strategy
in liposome-mediated gene therapy of bladder cancer is dependent on
sequence and order)

REFERENCE COUNT:

THERE ARE 43 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 22 OF 32 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1999:811109 HCAPLUS

43

DOCUMENT NUMBER:

132:69323

TITLE:

Prostate-associated antigen composition with chitosan metal chelate for the treatment of prostatic carcinoma

INVENTOR(S):

Seid, Christopher Allen; Singh, Gurpreet

PATENT ASSIGNEE(S):

Zonagen, Inc., USA PCT Int. Appl., 65 pp.

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO. DATE
wo 9965521	A1 19991223	WO 1999-US9592 19990430
W: AU, CA, RW: AT, BE,	CN, JP CH, CY, DE, DK,	ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
PT, SE US 2001014334	A1 20010816	US 1998-99017 19980617
US 6280742 AU 9936737	B2 20010828 A1 20000105 A1 20010404	AU 1999-36737 19990430 EP 1999-918940 19990430
EP 1087786 R: AT, BE,	DTZ FIG	TE NE CE MC DT
IE, FI JP 2002518345 IORITY APPLN. INFO	T2 20020625	JP 2000-554399 19990430 US 1998-99017 A 19980617

WO 1999-US9592 The present invention relates generally to materials and methods for redn. AΒ and/or alleviation of prostatic and prostatic-related (metatastic) carcinoma via the administration of compns. comprising a prostate-assocd. antigen and a chitosan-metal chelate.

158571-62-1, Lipofectamine IT

RL: BUU (Biological use, unclassified); BIOL (Biological study); USES

(prostate-assocd. antigen compn. with chitosan metal chelate for the treatment of prostatic carcinoma)

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

W 19990430

L11 ANSWER 23 OF 32 HCAPLUS COPYRIGHT 2002 ACS

2

ACCESSION NUMBER:

1999:679109 HCAPLUS

DOCUMENT NUMBER:

132:164839

TITLE:

Adjuvants for plasmid DNA vaccines

AUTHOR(S):

Norman, Jon; Hartikka, Jukka; Strauch, Pamela;

Manthorpe, Marston

CORPORATE SOURCE:

Vical Inc., San Diego, CA, USA

SOURCE:

Methods in Molecular Medicine (2000), 29, 185-196

CODEN: MMMEFN

PUBLISHER:

Humana Press Inc.

DOCUMENT TYPE:

Journal; General Review

English

A review with 38 refs. discussing the effects of the co-injection of LANGUAGE: bupivacaine (BP), polyvinyl pyrollidone (PVP), or DMRIE: DOPE cationic liposomes on plasmid DNA-mediated luciferase gene expression and antibody responses to influenza nucleoprotein (NP) antigen.

153312-64-2, DMRIE

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES

(DMRIE/DOPE liposomes contg.; adjuvants for plasmid DNA vaccines)

THERE ARE 38 CITED REFERENCES AVAILABLE FOR THIS 38 REFERENCE COUNT: RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 24 OF 32 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1998:795154 HCAPLUS

DOCUMENT NUMBER:

130:33989

TITLE:

Integrin-targeting vectors having transfection

activity

INVENTOR(S):

Hart, Stephen Lewis

PATENT ASSIGNEE(S):

Institute of Child Health, UK

SOURCE:

PCT Int. Appl., 70 pp.

DOCUMENT TYPE:

CODEN: PIXXD2

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.				KIND DATE			APPLICATION NO. DAY							_			
	WO	9854	 347		- -	 1	1998	1203		W	0 19	98 - GI	B157	7	1998	0529	~=	5.5
		W:	AL,	AM,	AT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CU,	CZ,	DE,
			DK.	EE.	ES.	FI,	GB,	GE,	GH,	GM,	GW,	HU,	ID,	IL,	IS,	JP,	ΚE,	KG,
			KP.	KR.	KZ.	LC.	LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,
			NO.	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,
			IIA.	UG.	US.	UZ.	VN,	YU,	ZW,	AM,	ΑZ,	ΒY,	KG,	KZ,	MD,	RU,	TU,	J.M
		RW:	GH.	GM.	KE.	LS.	MW,	SD,	SZ,	UG,	ZW,	ΑT,	BE,	CH,	CY,	DE,	DK,	ES,
		1000	FT.	FR.	GB,	GR.	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	CG,	CI,
			CM.	GA.	GN.	ML,	MR,	NE,	SN,	TD,	TG							
	וומ	9876		,	A	1	1998	1230	•	A	U 19	98-7	6673		1998	0529		
	EP	1003	898		A	1	2000	0531		E	P 19	98-9	2447	8	1998	0529		
	ы	R·	AT.	BE.	CH.	DE.	DK.	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
			IE,		,		•	•	·									
	.TP	2002			Т	2	2002	0122		J	P 19	98-5	5077	-	1998			
		2002					2002	0411		U	S 19	99-4	2465	6	1999	1129		
DDTO		Y APP				_				GB 1	997-	1111	5	Α	1997	0529		
EKTOI	. \	T 171 T		10	• •						998-				1998			
																n hi	ndin	~

A complex that comprises (1) a nucleic acid, (2) an integrin-binding AΒ component, for example, an integrin-binding peptide, (3) a polycationic nucleic acid-binding component, for example, oligolysine, and (4) a lipid component, for example, a cationic liposome, has transfection activity. Human neuroblastoma lines cells were transfected with a complex contg. lipofectin, the peptide K16-GACRRETAWACG with a nucleotide-binding domain and an .alpha.5.beta.1 integrin-binding domain, and retroviral vectors expressing interleukin-12 chains. Transfected cells secreted interleukin-12, demonstrating that the transection system is suitable for use in a vaccine for neuroblastoma and other cancers.

168479-03-6, DOSPA ΙT

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (integrin-targeting vectors having transfection activity)

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS 6 REFERENCE COUNT: RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 25 OF 32 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1998:789045 HCAPLUS

DOCUMENT NUMBER:

130:24103

TITLE:

An influenza enveloped DNA vaccine

INVENTOR(S):

Cusi, Maria Grazia; Gluck, Reinhard; Walti, Ernst

PATENT ASSIGNEE(S):

Schweiz. Serum- & Impfinstitut Bern, Switz.

PCT Int. Appl., 43 pp.

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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APPLICATION NO. DATE
                      KIND DATE
     PATENT NO.
                                                     _____
                                   _____
                                                     WO 1998-EP3050 19980522
     WO 9852603 .
                                   19981126
                           A2
                                   19990514
     WO 9852603
                           А3
          W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE,
               DK, EE, ES, FI, GB, GE, GH, GM, GW, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT,
                UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
          RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG
                                                                            19980522
                                                     AU 1998-79153
                            A1 19981211
     AU 9879153
                                                      EP 1998-929369 19980522
                            A2
                                   20000329
      EP 988052
           R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
                IE, FI
                                                                            19970523
                                                   EP 1997-108390
PRIORITY APPLN. INFO.:
                                                                            19980522
                                                   WO 1998-EP3050
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Described are virosomes comprising cationic lipids, biol. active influenza hemagglutinin protein or biol. active derivs. thereof and nucleic acids encoding antigens from pathogenic sources in their insides, preferably antigens from mumps virus wherein said antigens are derived from conserved external and internal proteins of said virus. Provided are virosomes which may advantageously be formulated as vaccines capable of inducing strong neutralizing antibody and cytotoxic T cell responses as well as protection to pathogenic sources such as a mumps virus. Furthermore, vaccines comprising recombinant DNA derived from DNA encoding conserved external and internal proteins from mumps virus are described. Mol. cloning of hemagglutinin gene, F gene, and nucleocapsid gene of mumps virus, N gene of respiratory syncytial virus, and S or Pre-S1 or Pre-S2 or S ORF gene of hepatitis B virus was described. Also described were prepn. of DOTAP-PC virosomes and DOTAP-PC-PE virosomes, incorporation of plasmids expressing mumps genes into DOTAP virosomes, humoral and cellular immune response to viral mumps-antigens induced by genetic immunization.

168479-03-6, DOSPA IT

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (virosomes comprising cationic lipids, influenza hemagglutinin, and antigen gene of pathogen as DNA vaccine for infectious diseases)

L11 ANSWER 26 OF 32 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1998:736678 HCAPLUS

DOCUMENT NUMBER:

130:91045

TITLE:

Direct gene transfer to the respiratory tract of mice

with pure plasmid and lipid-formulated DNA

AUTHOR(S):

McCluskie, Michael J.; Chu, Yongliang; Xia, Jiu-Lin; Jessee, Joel; Gebyehu, Gulilat; Davis, Heather L.

Loeb Research Institute, Ottawa, Can.

CORPORATE SOURCE:

SOURCE:

Antisense & Nucleic Acid Drug Development (1998),

8(5), 401-414

CODEN: ANADF5; ISSN: 1087-2906

Mary Ann Liebert, Inc. PUBLISHER:

DOCUMENT TYPE: LANGUAGE:

Journal English

Direct gene transfer into the respiratory system could be carried out for either therapeutic or immunization purposes. Here we demonstrate that AB cells in the lung can take up and express plasmid DNA encoding a luciferase reporter gene whether it is administered in naked form or formulated with cationic liposomes. Depending on the lipid used, the transfection efficiency with liposome-formulated DNA may be higher, the same as, or less than that with pure plasmid DNA. Tetramethyltetraalkylspermine analogs with alkyl groups of 16 or 18 carbons and DMRIE/cholesterol formulations proved particularly effective. Similar results for reporter gene expression in the lung were obtained whether the DNA (naked or lipid formulated) was administered by indirect, non-invasive intranasal delivery (inhaled or instilled) or by invasive, direct intratracheal delivery (injected or via a cannula). Reporter gene expression peaks around 4 days, then falls off dramatically by 9 days. The dose-response is linear, at least up to 100 .mu.g plasmid DNA, suggesting better transfection efficiencies might be realized if there was not a vol. limitation. For a given dose of DNA, the best results are obtained when the DNA is mixed with the min. amt. of lipid that can complex it completely. These results are discussed in the context of direct gene transfer for either gene therapy or delivery of a mucosal DNA vaccine.

153312-64-2, DMRIE ΙT

RL: BPR (Biological process); BSU (Biological study, unclassified); BUU (Biological use, unclassified); BIOL (Biological study); PROC (Process); USES (Uses)

(direct gene transfer to respiratory tract of mice with pure plasmid and lipid-formulated DNA)

REFERENCE COUNT:

THERE ARE 71 CITED REFERENCES AVAILABLE FOR THIS 71 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 27 OF 32 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1998:548562 HCAPLUS

DOCUMENT NUMBER:

129:193718

TITLE:

Formulation of stabilized cationic transfection agents

complexed with nucleic acid particles

INVENTOR(S):

Crouzet, Joel; Pitard, Bruno Rhone-Poulenc Rorer S.A., Fr.

PATENT ASSIGNEE(S):

PCT Int. Appl., 52 pp.

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

LANGUAGE:

Patent French

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.			KIND	D DATE		APPLICATION NO. DATE										
WO	T 0	TD	BA, BB,	τư	BR,	T.T.	CN,	CU, MG.	MK.	EE, MN,	GE, MX,	NO,	HU, NZ,	,	110,	
	FR.	GM, GB,	TJ, TM KE, LS, GR, IE, ML, MR,	IT, NE,	LU, SN,	MC, TD,	NL, TG	PT,	SE,	Br,	BU,	DE, CF,	cu,	C1,	FI, CM,	
	2759298		A1 B1	1998			F.	R 19	97-1	46/		1991	0210			
	2759298 9862987		A1	1998	0826		A	U 19	98-6	2987		1998	0206			
AU 737720 BR 9807563 EP 1007097		B2 A A1	2001 2000 2000	BR 1998-7563 EP 1998-906986							L9980206 L9980206					

20011017 В1 EP 1007097 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, PT, IE, SI, FI JP 1998-533881 19980206 T2 20010807 JP 2001511171 19980206 AT 1998-906986 20011115 AT 206932 E 19980206 ES 1998-906986 Т3 20020401 ES 2166146 ZA 1998-1034 19980209 19980811 ZA 9801034 Α NO 1999-3825 19990809 Α 19990809 NO 9903825 FR 1997-1467 A 19970210 PRIORITY APPLN. INFO.: WO 1998-FR222 W 19980206

MARPAT 129:193718 OTHER SOURCE(S):

The invention concerns a compn. contg. stabilized particles of cationic transfection agent(s)/nucleic acid complexes characterized in that it includes besides said transfection agent and nucleic acid at least a non-ionic surfactant in sufficient amt. for preventing the aggregation of the particles in course of time. In a preferred embodiment, the surfactant is a polyoxyalkylene or a deriv. thereof.

158571-62-1, Lipofectamine

RL: DEV (Device component use); PEP (Physical, engineering or chemical process); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)

(formulation of stabilized cationic transfection agents complexed with nucleic acid particles)

L11 ANSWER 28 OF 32 HCAPLUS COPYRIGHT 2002 ACS

1998:249878 HCAPLUS ACCESSION NUMBER:

129:12373 DOCUMENT NUMBER:

Transfection of primary tumor cells and tumor cell TITLE:

lines with plasmid DNA/lipid complexes

Stopeck, Alison T.; Hersh, Evan M.; Brailey, AUTHOR(S):

Jacqueline L.; Clark, Paul R.; Norman, Jon; Parker,

Suezanne E.

Arizona Cancer Center, Tucson, AZ, 85724-5024, USA CORPORATE SOURCE:

Cancer Gene Therapy (1998), 5(2), 119-126 SOURCE:

CODEN: CGTHEG; ISSN: 0929-1903

Appleton & Lange

PUBLISHER:

Journal DOCUMENT TYPE: English LANGUAGE:

Cancer vaccines that utilize genetically modified tumor cells require gene transfer methods capable of producing immunostimulatory doses of transgenes from fresh or short-term cultures of human tumor cells. Our studies optimize in vitro transfection of primary tumor cells using cationic lipids and a plasmid encoding the gene for human interleukin-2 (IL-2). Established tumor cell lines produced 10- to 100-fold more IL-2 than did fresh or short-term tumor cultures as measured by enzyme-linked immunoabsorbent anal. Importantly, transfection of primary tumor cells produced immunostimulatory levels of IL-2 as detd. by increased thymidine incorporation by autologous peripheral blood mononuclear cells and lymphokine-activated killer cell activity. IL-2 secretion by tumor cells persisted for at least 30 days post-transfection and was unaffected by freeze thawing or irradn. to 8000 rads. Multiple solid tumor types were successfully transfected, but normal blood mononuclear cells and leukemic blasts were resistant to transfection. Enzyme-linked immunoabsorbent anal. of the amt. of IL-2 secreted into the medium by transfected tumor cells correlated with the percentage of tumor cells expressing intracellular IL-2 as measured by flow cytometry. Plasmids utilizing a cytomegalovirus promoter yielded superior transfection efficiencies compared with plasmids contg. a Rous sarcoma virus promoter. These results suggest that a clin. vaccine trial using autologous tumor cells genetically modified to secrete IL-2 is feasible in patients

with solid tumors. 153312-64-2, DMRIE

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (primary tumor cell and tumor cell line transfection with IL-2-encoding plasmid DNA/cationic lipid complexes)

L11 ANSWER 29 OF 32 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1998:180751 HCAPLUS

DOCUMENT NUMBER:

128:248559

TITLE:

ΙT

Cationic liposomes with entrapped polynucleotides for

use as gene vaccines Gregoriadis, Gregory

INVENTOR(S): PATENT ASSIGNEE(S):

School of Pharmacy, UK; Gregoriadis, Gregory

SOURCE:

PCT Int. Appl., 51 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO. DATE
WO 9810748 W: AL, AM, DK, EE, KZ, LC, PL, PT,	A1 19980319 AT, AU, AZ, BA, BB, ES, FI, GB, GE, GH, LK, LR, LS, LT, LU, RO, RU, SD, SE, SG, VN, YU, ZW, AM, AZ,	WO 1997-GB2490 19970915 BG, BR, BY, CA, CH, CN, CU, CZ, DE, HU, ID, IL, IS, JP, KE, KG, KP, KR, LV, MD, MG, MK, MN, MW, MX, NO, NZ, SI, SK, SL, TJ, TM, TR, TT, UA, UG, BY, KG, KZ, MD, RU, TJ, TM
GB, GR, GN, MI,	IE, IT, LU, MC, NL, MR, NE, SN, TD, TG	ZW, AT, BE, CH, DE, DK, ES, FI, FR, PT, SE, BF, BJ, CF, CG, CI, CM, GA,
AU 9742154	A1 19980402 B2 20010111	
R: AT, BE, CN 1237102 JP 2001502299	CH, DE, DK, ES, FR, A 19991201 T2 20010220 A 20000626	KR 1999-702103 19990312
PRIORITY APPLN. INFO		GB 1996-19172 A 19960913 GB 1996-25917 A 19961213 GB 1997-13994 A 19970701 WO 1997-GB2490 W 19970915

MARPAT 128:248559 OTHER SOURCE(S):

Cationic liposomes with entrapped polynucleotide in the intravesicular space are described. The liposomes include cationic components such as cationic lipids such as DOTAP. Preferably the method of forming liposomes uses the dehydration-rehydration method in the presence of the polynucleotide. The polynucleotide preferably operatively encodes an antigen capable of eliciting a desired immune response, i.e., is a gene vaccine.

158571-62-1, Lipofectamine ΙT

RL: DEV (Device component use); PEP (Physical, engineering or chemical process); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)

(cationic liposomes with entrapped polynucleotides for use as gene vaccines)

L11 ANSWER 30 OF 32 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1997:473805 HCAPLUS

DOCUMENT NUMBER:

127:175118

ANGELL 09 / 760574

TITLE:

Development of improved vectors for DNA-based immunization and other gene therapy applications Norman, Jon A.; Hobart, Peter; Manthorpe, Marston;

AUTHOR(S):

Felgner, Phil; Wheeler, Carl

CORPORATE SOURCE:

Vical Inc., San Diego, CA, 92121, USA

SOURCE:

Vaccine (1997), 15(8), 801-803 CODEN: VACCDE; ISSN: 0264-410X

PUBLISHER: DOCUMENT TYPE: Elsevier Journal

English LANGUAGE:

Optimizing gene expression and delivery are necessary steps in the prodn. AB of vectors for DNA-based immunization as well as for other gene therapy applications. A mouse muscle/reporter gene assay system was used to systematically improve a plasmid DNA vector. The optimized vector VR1255 contained: (1) CMV promoter and enhancer; (2) CMV IE Intron A; (3) kanamycin resistance gene; (4) deleted SV40 origin of replication; (5) optimized lux coding region; and (6) a minimal synthetic terminator from the rabbit beta globin gene, mRBG. The vector VR1255 expressed 137 times greater than an earlier prototype RSV-based vector. For plasmid vector delivery into nonmuscle tissues, a recently synthesized cationic lipid, GAP-DLRIE, was found to greatly enhance the uptake and expression of plasmid DNA by 100-fold when instilled into the mouse lung. The time-course of CAT expression with GAP-DLRIE indicated that peak expression occurs 2-5 days after intranasal administration and expression diminished to about one-third the peak value by day 21. This cationic lipid may be useful for immunization by pulmonary and perhaps other nonmuscle routes.

182919-20-6P ΙT

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(development of improved vectors for DNA-based immunization and other

gene therapy applications)

L11 ANSWER 31 OF 32 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1997:429591 HCAPLUS

DOCUMENT NUMBER:

127:49213

TITLE:

Novel non-pyrogenic bacterial strains and use of the

same

INVENTOR(S):

Hone, David M.; Powell, Robert J.

PATENT ASSIGNEE(S):

University of Maryland At Baltimore, USA; Hone, David

M.; Powell, Robert J.

SOURCE:

PCT Int. Appl., 124 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT N	KIND DATE				A.	PPLI	CATI	ο. :	DATE								
					_												
WO 9718837			A1 19970529					WO 1996-US19875 19961122									
W:	AL.	AM.	AT,	AU,	AZ,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CZ,	DE,	DK,	EE,	
	ES,	FI,	GB,	GE,	HU,	IL,	IS,	JP,	ΚE,	KG,	ΚP,	KR,	ΚZ,	LK,	LR,	LS,	
	LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	ΝZ,	PL,	PT,	RO,	RU,	SD,	
	SE,	SG,	SI,	SK,	ТJ,	TM,	TR,	TT,	UA,	UG,	US,	UZ,	VN,	AM,	ΑZ,	BY,	
	KG,	KZ,	MD,	RU,	ТJ,	TM											
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	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ΒJ,	CF,	CG,	CI,	CM,	GA,	GN,	ML,	
	MR,	NE,	SN,	TD,	TG												

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AU 1997-22784
                                                            19961122
                      Α1
                            19970611
    AU 9722784
                                           EP 1996-945937
                                                            19961122
                            19980520
    EP 841941
                      Α1
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, FI
                            19991207
                                           US 1997-802371
                                                            19970219
     US 5997881
                                        US 1995-7478P
                                                         P 19951122
PRIORITY APPLN. INFO.:
                                        WO 1996-US19875 W 19961122
```

AB The present invention provides gram-neg. bacterial strains that produce substantially pure non-pyrogenic lipopolysaccharide or lipid A. The present invention also relates to a use of said strains for the prepn. of non-pyrogenic DNA and use of the same for introducing endogenous or foreign genes into animal cells or animal tissue. Further, the present invention relates to a use of said strains for the prepn. of non-pyrogenic bacterial proteins and polysaccharides antigens for use as vaccines.

IT 168479-03-6

RL: MOA (Modifier or additive use); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(non-pyrogenic bacterial strains producing non-pyrogenic lipid A for delivery vaccine genes or DNA into animal cell or tissue)

L11 ANSWER 32 OF 32 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1996:590320 HCAPLUS

DOCUMENT NUMBER: 125:212664

TITLE: Combined therapeutic treatment of hyperproliferative

diseases using oncogenic cell-signaling

pathway-inhibiting nucleic acids and anticancer agents

INVENTOR(S): Tocque, Bruno

PATENT ASSIGNEE(S): Rhone-Poulenc Rorer S.A., Fr.

SOURCE: PCT Int. Appl., 35 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: French

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	PATENT NO.				ND	DATE			A	PPLI	CATI	ON NO). 	DATE						
WO	9622101			A1			 A1			0725		W	0 19	96-F	R56	1996	0112			
	W:	AM,	AU,	BB.	BG,	BR,	BY,	CA,	CN,	CZ,	EE,	FΙ,	GΕ,	HU,	IS,	JP,	KG,			
		KP,	KR,	ΚZ,	LK,	LR,	LT,	LV,	MD,	MG,	MN,	MX,	NO,	NZ,	PL,	RO,	RU,			
		SG.	SI,	SK,	TJ,	TM,	TT,	UA,	UG,	US,	UZ,	VN								
	RW:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙE,	ΙΤ,	LU,	MC,	NL,	PT,	SE,			
		BF.	ВJ.	CF,	CG,	CI,	CM,	GΑ,	GN,	ML,	MR,	ΝE,	SN,	TD,	TG					
FR	2729295 2729295 2209771			Α	1	19960719			FR 1995-436					1995	0117					
FR				В	1	1997	0228													
CA				A	A	1996	0725		CA 1996-2209771					1996	0112					
ΔIJ				A1		19960807			A	AU 1996-45429				19960112						
AU	7163	64		В	2	2000	0224													
EΡ	8003	99		Α	1	1997	1015		Ε	P 19	96-9	0138	7	1996	0112			0 T		
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	PT,	IE,	SI		
BR	9606	969		Α		1997	1104		В	R 19	996-6	969	_	1996	0112					
JP	1051	2559		T	2	1998	1202		J	P 19	996-5	2207	8	1996	0112					
NO	9703	197		A		1997	0709		N	0 19	97-3	197		1997	0709					
FI	9703	023		A		1997	0716		FI 1997-3023 US 1997-875222 US 2001-816144					1997	0715					
US	6262	032		В	1	2001	0717							1997	0717					
US	2001	0213	95	A	1	2001	0913		0	S 20	001-8	1614	4	200I	0326					
CORIT	ORITY APPLN. INFO			.:					FR I	995-	-436		A	1995	0117					
									WO I	996-	-FR56		W 70.1	1996	0112					
							US I	99/-	-8/52		ΑI	1997	0/1/							

OTHER SOURCE(S): MARPAT 125:212664

AB Hyperproliferative diseases are treated with a medicinal combination of .gtoreq.1 nucleic acids that at least partially inhibit oncogenic cell signaling pathways, and a therapeutic anticancer agent. The nucleic acid is e.g. a DNA coding for a tumor suppressor protein. The anticancer agent may be a taxoid, vinca alkaloid, etc.

IT 158571-62-1

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (hyperproliferative disease combined therapeutic treatment with oncogenic cell-signaling pathway-inhibiting nucleic acids and anticancer agents)

=> sel hit rn E1 THROUGH E19 ASSIGNED

=> file reg FILE 'REGISTRY' ENTERED AT 14:56:02 ON 16 AUG 2002 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2002 American Chemical Society (ACS)

STRUCTURE FILE UPDATES: 15 AUG 2002 HIGHEST RN 444046-42-8 DICTIONARY FILE UPDATES: 15 AUG 2002 HIGHEST RN 444046-42-8

TSCA INFORMATION NOW CURRENT THROUGH MAY 20, 2002

Please note that search-term pricing does apply when conducting ${\tt SmartSELECT}$ searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Calculated physical property data is now available. See HELP PROPERTIES for more information. See STNote 27, Searching Properties in the CAS Registry File, for complete details: http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf

=> s e1-e19

1 153312-64-2/BI (153312-64-2/RN) 1 158571-62-1/BI (158571-62-1/RN) 1 168479-03-6/BI (168479-03-6/RN) 1 208040-06-6/BI (208040-06-6/RN) 1 299207-54-8/BI (299207-54-8/RN) 1 182919-20-6/BI (182919-20-6/RN) 1 370108-99-9/BI (370108-99-9/RN) 1 153312-60-8/BI (153312-60-8/RN) 1 154486-25-6/BI (154486-25-6/RN) 1 188949-12-4/BI (188949-12-4/RN) 1 189203-05-2/BI (189203-05-2/RN)

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                 (199171-54-5/RN)
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                 (282533-23-7/RN)
             1 282533-25-9/BI
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                 (299207-55-9/RN)
             1 310445-42-2/BI
                 (310445-42-2/RN)
             1 310445-43-3/BI
                 (310445-43-3/RN)
             1 310445-44-4/BI
                 (310445-44-4/RN)
             1 370108-98-8/BI
                 (370108-98-8/RN)
            19 (153312-64-2/BI OR 158571-62-1/BI OR 168479-03-6/BI OR 208040-06
L12
               -6/BI OR 299207-54-8/BI OR 182919-20-6/BI OR 370108-99-9/BI OR
               153312-60-8/BI OR 154486-25-6/BI OR 188949-12-4/BI OR 189203-05-
               2/BI OR 199171-54-5/BI OR 282533-23-7/BI OR 282533-25-9/BI OR
               299207-55-9/BI OR 310445-42-2/BI OR 310445-43-3/BI OR 310445-44-
               4/BI OR 370108-98-8/BI)
=> d ide can 112 tot
L12 ANSWER 1 OF 19 REGISTRY COPYRIGHT 2002 ACS
     370108-99-9 REGISTRY
RN
     1-Propanaminium, N-(3-aminopropyl)-N,N-dimethyl-2,3-bis[(9Z)-9-
CN
     tetradecenyloxy]-, bromide, mixt. with (1R)-1-[[[(2-
     aminoethoxy)hydroxyphosphinyl]oxy]methyl]-1,2-ethanediyl
     bis(3,7,11,15-tetramethylhexadecanoate) (1:1) (9CI) (CA INDEX NAME)
OTHER CA INDEX NAMES:
     Hexadecanoic acid, 3,7,11,15-tetramethyl-, (1R)-1-[[[(2-
     aminoethoxy)hydroxyphosphinyl]oxy]methyl]-1,2-ethanediyl ester, mixt.
     contg. (9CI)
OTHER NAMES:
     Vaxfectin
CN
     STEREOSEARCH
FS
     C45 H90 N O8 P . C36 H73 N2 O2 . Br
MF
CI
     MXS
SR
                  CA, CAPLUS, TOXCENTER
LC
     STN Files:
     CM
          1
     CRN
          370108-98-8
         C36 H73 N2 O2 . Br
     CMF
```

Double bond geometry as shown.

$$n-Bu$$
 Z
 $CH_2)_8$
 Me
 Me
 $CH_2)_3$
 NH_2
 NH_2

● Br-

CM 2

CRN 201036-16-0 CMF C45 H90 N O8 P

Absolute stereochemistry.

PAGE 1-A

$$H_2N$$
 H_2N
 $H_$

PAGE 1-B

$$\begin{array}{c} \text{Me} & \text{Me} \\ \text{(CH2)} & \text{3} \end{array}$$
 (CH2) $\begin{array}{c} \text{CHMe2} \\ \text{(CH2)} & \text{3} \end{array}$

2 REFERENCES IN FILE CA (1967 TO DATE)
2 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 136:198465

REFERENCE 2: 135:330213

L12 ANSWER 2 OF 19 REGISTRY COPYRIGHT 2002 ACS

RN **370108-98-8** REGISTRY

CN 1-Propanaminium, N-(3-aminopropyl)-N,N-dimethyl-2,3-bis[(9Z)-9-tetradecenyloxy]-, bromide (9CI) (CA INDEX NAME)
OTHER NAMES:

CN VC 1052

FS STEREOSEARCH

MF C36 H73 N2 O2 . Br

CI COM

SR CA

,

LC STN Files: CA, CAPLUS, TOXCENTER

Double bond geometry as shown.

$$n-Bu$$

$$Z$$
 $(CH_2)_8$

$$N$$

$$CH_2)_8$$

$$N$$

$$+$$

$$CH_2)_3$$

$$NH_2$$

$$NH_2$$

• Br-

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 135:330213

L12 ANSWER 3 OF 19 REGISTRY COPYRIGHT 2002 ACS

RN **310445-44-4** REGISTRY

CN 16-Oxa-4,7-diaza-12-azoniatriacontan-1-aminium, N-[2,3-bis(tetradecyloxy)propyl]-6-(1H-indol-3-ylmethyl)-N,N,12,12-tetramethyl-5,8-dioxo-14-(tetradecyloxy)-, dibromide, (6S)- (9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C84 H161 N5-O6 . 2 Br

SR CA

LC STN Files: CA, CAPLUS

Absolute stereochemistry.

●2 Br-

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 134:21482

L12 ANSWER 4 OF 19 REGISTRY COPYRIGHT 2002 ACS

RN **310445-43-3** REGISTRY

CN 4,6,11,13-Tetraazahexadecane-1,16-diaminium, N,N'-bis[2,3-bis(tetradecyloxy)propyl]-N,N,N',N'-tetramethyl-5,12-dioxo-(9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C78 H162 N6 O6

SR CA

LC STN Files: CA, CAPLUS

PAGE 1-B

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 134:21482

L12 ANSWER 5 OF 19 REGISTRY COPYRIGHT 2002 ACS

RN **310445-42-2** REGISTRY

CN 4,6,13,15-Tetraazaoctadecane-1,18-diaminium, N,N'-bis[2,3-bis(tetradecyloxy)propyl]-N,N,N',N'-tetramethyl-5,14-dioxo-(9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C80 H166 N6 O6

SR CA

LC STN Files: CA, CAPLUS

PAGE 1-B

1 REFERENCES IN FILE CA (1967 TO DATE)
1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 134:21482

L12 ANSWER 6 OF 19 REGISTRY COPYRIGHT 2002 ACS

RN 299207-55-9 REGISTRY

CN 1-Propanaminium, N-(2-aminoethyl)-2,3-bis(hexadecyloxy)-N,N-dimethyl-, bromide (9CI) (CA INDEX NAME)

OTHER NAMES:

CN GAP-DPRIE

MF C39 H83 N2 O2 . Br

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER

● Br -

1 REFERENCES IN FILE CA (1967 TO DATE)
1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 133:280556

L12 ANSWER 7 OF 19 REGISTRY COPYRIGHT 2002 ACS

RN 299207-54-8 REGISTRY

CN 1-Propanaminium, N-(2-aminoethyl)-N,N-dimethyl-2,3-bis[(9Z)-9-tetradecenyloxy]-, bromide (9CI) (CA INDEX NAME)

OTHER NAMES:

CN GAP-DMORIE

FS STEREOSEARCH

MF C35 H71 N2 O2 . Br

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

Double bond geometry as shown.

$$rac{D}{Z}$$
 $(CH_2)_8$ O Me Me NH_2 $n-Bu$

• Br-

3 REFERENCES IN FILE CA (1967 TO DATE)
3 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 135:348865

REFERENCE 2: 134:161880

REFERENCE 3: 133:280556

L12 ANSWER 8 OF 19 REGISTRY COPYRIGHT 2002 ACS

RN 282533-25-9 REGISTRY

CN 1-Propanaminium, N-(3-aminopropyl)-N,N-dimethyl-2,3-bis(tetradecyloxy)-, bromide (9CI) (CA INDEX NAME)

MF C36 H77 N2 O2 . Br

SR CA

LC STN Files: CA, CAPLUS

CRN (191980-83-3)

• Br-

3 REFERENCES IN FILE CA (1967 TO DATE)

3 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 135:142076

REFERENCE 2: 134:21482

REFERENCE 3: 133:103732

L12 ANSWER 9 OF 19 REGISTRY COPYRIGHT 2002 ACS

RN 282533-23-7 REGISTRY

CN 1-Propanaminium, N-[2-[[2,5-bis[(3-aminopropyl)amino]-1-oxopentyl]amino]ethyl]-N,N-dimethyl-2,3-bis[(9Z)-9-octadecenyloxy]-, chloride, tetrahydrochloride (9CI) (CA INDEX NAME)

OTHER NAMES:

CN DOSPA

FS STEREOSEARCH

MF C54 H111 N6 O3 . 4 Cl H . Cl

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER

Double bond geometry as shown.

$$(CH_2)_3$$
 $(CH_2)_3$
 $(CH_2)_3$

● Cl⁻

• 4 HCl

PAGE 1-B

(CH₂)7Me (CH₂)7Me

4 REFERENCES IN FILE CA (1967 TO DATE)

4 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 136:314910

REFERENCE 2: 133:340208

REFERENCE 3: 133:280556

REFERENCE 4: 133:103732

L12 ANSWER 10 OF 19 REGISTRY COPYRIGHT 2002 ACS

RN **208040-06-6** REGISTRY

CN 1-Propanaminium, N-(2-aminoethyl)-2,3-bis(dodecyloxy)-N,N-dimethyl-, bromide (9CI) (CA INDEX NAME)

OTHER NAMES:

CN GAP-DLRIE

MF C31 H67 N2 O2 . Br

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

• Br-

5 REFERENCES IN FILE CA (1967 TO DATE)

5 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 135:348865

REFERENCE 2: 134:161880

REFERENCE 3: 133:340208

REFERENCE 4: 133:280556

REFERENCE 5: 129:32388

L12 ANSWER 11 OF 19 REGISTRY COPYRIGHT 2002 ACS

RN 199171-54-5 REGISTRY

CN 1-Propanaminium, 2,3-bis(dodecyloxy)-N-(2-hydroxyethyl)-N,N-dimethyl-, bromide (9CI) (CA INDEX NAME)

OTHER NAMES:

CN DLRIE

MF C31 H66 N O3 . Br

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

• Br-

7 REFERENCES IN FILE CA (1967 TO DATE)

7 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 135:142076

REFERENCE 2: 133:340208

REFERENCE 3: 133:280556

REFERENCE 4: 133:103732

REFERENCE 5: 132:298688

REFERENCE 6: 131:14825

REFERENCE 7: 128:16429

L12 ANSWER 12 OF 19 REGISTRY COPYRIGHT 2002 ACS

RN 189203-05-2 REGISTRY

CN Cholest-5-en-3-ol (3.beta.)-, mixt. with N-(2-hydroxyethyl)-N,N-dimethyl-2,3-bis(tetradecyloxy)-1-propanaminium bromide (9CI) (CA INDEX NAME)
OTHER CA INDEX NAMES:

CN 1-Propanaminium, N-(2-hydroxyethyl)-N, N-dimethyl-2, 3-bis(tetradecyloxy)-, bromide, mixt. contg. (9CI)

OTHER NAMES:

CN Cholesterol mixt. with DMRIE

CN DMRIE-C

CN DMRIE-cholesterol mixt.

FS STEREOSEARCH

MF C35 H74 N O3 . C27 H46 O . Br

CI MXS

SR CA

LC STN Files: BIOSIS, CA, CAPLUS, TOXCENTER, USPATFULL

CM 1

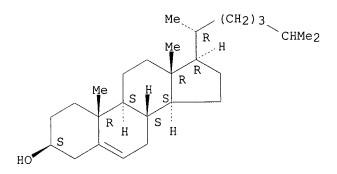
CRN 153312-64-2 (191980-81-1) CMF C35 H74 N O3 . Br

• Br-

CM 2

CRN 57-88-5 CMF C27 H46 O

Absolute stereochemistry.



30 REFERENCES IN FILE CA (1967 TO DATE)
30 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 137:72660

REFERENCE 2: 136:350922

REFERENCE 3: 136:227908

REFERENCE 4: 136:172607

REFERENCE 5: 136:42689

REFERENCE 6: 136:328

REFERENCE 7: 135:362424

REFERENCE 8: 135:262092

REFERENCE 9: 135:41420

REFERENCE 10: 135:14107

L12 ANSWER 13 OF 19 REGISTRY COPYRIGHT 2002 ACS

RN 188949-12-4 REGISTRY

CN 1-Propanaminium, N-(2-hydroxyethyl)-N,N-dimethyl-2,3-bis[(9Z)-9-tetradecenyloxy]-, bromide (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 1-Propanaminium, N-(2-hydroxyethyl)-N,N-dimethyl-2,3-bis(9-tetradecenyloxy)-, bromide, (Z,Z)-

OTHER NAMES:

CN DMORIE

FS STEREOSEARCH

MF C35 H70 N O3 . Br

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER

Double bond geometry as shown.

$$C$$
 $CH_2)_8$ C Me Me Me N O N

● Br-

2 REFERENCES IN FILE CA (1967 TO DATE)
2 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 133:280556

REFERENCE 2: 126:282608

L12 ANSWER 14 OF 19 REGISTRY COPYRIGHT 2002 ACS

RN 182919-20-6 REGISTRY

CN 1-Propanaminium, N-(3-aminopropyl)-2,3-bis(dodecyloxy)-N,N-dimethyl-, bromide (9CI) (CA INDEX NAME)

MF C32 H69 N2 O2 . Br

CI COM

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER

CRN (191980-99-1)

• Br-

11 REFERENCES IN FILE CA (1967 TO DATE)

11 REFERÈNCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 135:335117

REFERENCE 2: 135:142076

REFERENCE 3: 133:291661

REFERENCE 4: 133:103732

REFERENCE 5: 131:18016

REFERENCE 6: 131:14825

REFERENCE 7: 129:99906

REFERENCE 8: 127:175118

REFERENCE 9: 127:103864

REFERENCE 10: 125:338808

L12 ANSWER 15 OF 19 REGISTRY COPYRIGHT 2002 ACS

RN **168479-03-6** REGISTRY

CN 1-Propanaminium, N-[2-[[2,5-bis[(3-aminopropyl)amino]-1-

oxopentyl] amino] ethyl]-N, N-dimethyl-2, 3-bis[[(9Z)-1-oxo-9-octadecenyl]oxy]-

, salt with trifluoroacetic acid (1:1) (9CI) (CA INDEX NAME)

OTHER NAMES:

CN 2,3-Dioleoyloxy-N-[2-(sperminecarboxamido)ethyl]-N,N-dimethyl-1-propanaminium trifluoroacetate

CN DOSPA

FS STEREOSEARCH

DR 163046-76-2

MF C54 H107 N6 O5 . C2 F3 O2

CI COM

SR CA

LC STN Files: BIOBUSINESS, CA, CAPLUS, TOXCENTER, USPATFULL

CM 1

CRN 168479-02-5

CMF C54 H107 N6 O5

Double bond geometry as shown.

PAGE 1-A

$$(CH_2)_3$$
 $(CH_2)_3$
 $(CH_2)_3$
 $(CH_2)_3$
 $(CH_2)_3$
 $(CH_2)_3$
 $(CH_2)_3$
 $(CH_2)_3$
 $(CH_2)_3$
 $(CH_2)_3$
 $(CH_2)_3$

PAGE 1-B

CM 2

CRN 14477-72-6 CMF C2 F3 O2

65 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

65 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 137:83640

REFERENCE 2: 137:57284

REFERENCE 3: 136:336176

REFERENCE 4: 136:314966

REFERENCE 5: 136:307351

REFERENCE 6: 136:195950

REFERENCE 7: 136:123632

REFERENCE 8: 136:32635

REFERENCE 9: 136:32634

REFERENCE 10: 136:97

L12 ANSWER 16 OF 19 REGISTRY COPYRIGHT 2002 ACS

RN 158571-62-1 REGISTRY

CN 1-Propanaminium, N-[3-[[3-[[4-[(3-aminopropyl)amino]butyl]amino]propyl]amino]-3-oxopropyl]-N, N-dimethyl-2, 3-bis[[(9Z)-1-oxo-9-octadecenyl]oxy]-, salt with trifluoroacetic acid (1:1), mixt. with 1-[[[(2-aminoethoxy)hydroxyphosphinyl]oxy]methyl]-1, 2-ethanediyl

di-(9Z)-9-octadecenoate (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

1-Propanaminium, N-[3-[[3-[[4-[(3-aminopropyl)amino]butyl]amino]propyl]ami no]-3-oxopropyl]-N, N-dimethyl-2, 3-bis[(1-oxo-9-octadecenyl)oxy]-, (Z,Z)-, salt with trifluoroacetic acid (1:1), mixt. with (Z,Z)-1-[[[(2-1)]]aminoethoxy)hydroxyphosphinyl]oxy]methyl]-1,2-ethanediyl di-9-octadecenoate

9-Octadecenoic acid (9Z)-, 1-[[[(2-aminoethoxy)hydroxyphosphinyl]oxy]methyCN 1]-1,2-ethanediyl ester, mixt. contg. (9CI)

9-Octadecenoic acid (Z)-, 2-deoxy-2-[(1-oxododecyl)amino]-, CN 1-[[[(2-aminoethoxy)hydroxyphosphinyl]oxy]methyl]-1,2-ethanediyl ester, mixt. contg.

OTHER NAMES:

CN LipofectAMINE

FS STEREOSEARCH

C54 H106 N5 O5 . C41 H78 N O8 P . C2 F3 O2 MF

CI MXS

SR CA

LC AGRICOLA, BIOSIS, BIOTECHNO, CA, CAPLUS, CIN, EMBASE, IPA, STN Files: TOXCENTER, USPAT2, USPATFULL

CM 1

2462-63-7 CRN C41 H78 N O8 P

Double bond geometry as shown.

PAGE 1-A

$$H_2N$$
 H_2N
 $H_$

PAGE 1-B

__ Me

CM 2

185097-43-2 CRN C54 H106 N5 O5 . C2 F3 O2 CMF

> CM 3

CRN 181508-68-9 CMF C54 H106 N5 O5 Double bond geometry as shown.

$$H_2N$$
 (CH₂) 3 (CH₂) 4 (CH₂) 3 N Me Me Me O (CH₂) 7 (CH₂) 7

PAGE 1-B

CM 4

CRN 14477-72-6 CMF C2 F3 O2

221 REFERENCES IN FILE CA (1967 TO DATE)
8 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
221 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 137:92454

REFERENCE 2: 137:83524

REFERENCE 3: 137:76688

REFERENCE 4: 137:72673

REFERENCE 5: 137:68056

REFERENCE 6: 137:58112

REFERENCE 7: 137:57284

8: 137:42288 REFERENCE

9: 137:32101 REFERENCE

REFERENCE 10: 137:1159

L12 ANSWER 17 OF 19 REGISTRY COPYRIGHT 2002 ACS

154486-25-6 REGISTRY RN

1-Propanaminium, N-(2-aminoethyl)-N, N-dimethyl-2, 3-bis(tetradecyloxy)-, bromide (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

1-Propanaminium, N-(2-aminoethyl)-N, N-dimethyl-2, 3-bis(tetradecyloxy)-, bromide, (.+-.)-

OTHER NAMES:

GAP-DMRIE CN

C35 H75 N2 O2 . Br MF

CAS Registry Services SR

STN Files: CA, CAPLUS, TOXCENTER, USPATFULL LC

CRN (191980-79-7)

● Br-

- 5 REFERENCES IN FILE CA (1967 TO DATE)
- 5 REFERENCES IN FILE CAPLUS (1967 TO DATE)

1: 133:280556 REFERENCE

2: 133:103732 REFERENCE

3: 131:18016 REFERENCE

4: 129:36461 REFERENCE

5: 124:279113 REFERENCE

L12 ANSWER 18 OF 19 REGISTRY COPYRIGHT 2002 ACS

153312-64-2 REGISTRY

1-Propanaminium, N-(2-hydroxyethyl)-N,N-dimethyl-2,3-bis(tetradecyloxy)-, bromide (9CI) (CA INDEX NAME)

OTHER NAMES:

DMRIE CN

- N-[1-(2,3-Ditetradecyloxy)propyl]-N, N-dimethyl-N-hydroxyethylammoniumCN bromide
- 146659-77-0 DR
- C35 H74 N O3 . Br ΜF
- COM CI
- SR CA
- BIOSIS, CA, CANCERLIT, CAPLUS, IPA, MEDLINE, TOXCENTER, LC STN Files: USPATFULL

CRN (191980-81-1)

• Br-

110 REFERENCES IN FILE CA (1967 TO DATE) 6 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

110 REFERENCES IN FILE CAPLUS (1967 TO DATE)

1: 136:390857 REFERENCE

2: 136:336176 REFERENCE

3: 136:268001 REFERENCE

4: 136:195950 REFERENCE

5: 136:123632 REFERENCE

6: 136:107481 REFERENCE

7: 136:74555 REFERENCE

8: 136:58784 REFERENCE

9: 136:58673 REFERENCE

REFERENCE 10: 135:370419

L12 ANSWER 19 OF 19 REGISTRY COPYRIGHT 2002 ACS

153312-60-8 REGISTRY

1-Propanaminium, N-(2-hydroxyethyl)-N,N-dimethyl-2,3-bis[(9Z)-9octadecenyloxy]-, bromide (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

1-Propanaminium, N-(2-hydroxyethyl)-N,N-dimethyl-2,3-bis(9-octadecenyloxy)-, bromide, (Z,Z)-

OTHER NAMES:

CN DORIE

STEREOSEARCH FS

C43 H86 N O3 . Br MF

SR

CA, CAPLUS, TOXCENTER, USPATFULL LC STN Files:

CRN (153985-18-3)

Double bond geometry as shown.

Me (CH₂) 7
$$Z$$
 (CH₂) 8 Z (CH₂) 7 Me Me Me

• Br-

4 REFERENCES IN FILE CA (1967 TO DATE)

4 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 133:355219

REFERENCE 2: 133:280556

REFERENCE 3: 133:103732

REFERENCE 4: 120:153003